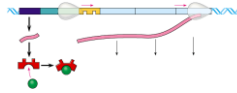




Control of Prokaryotic (Bacterial) Genes



Bacterial metabolism

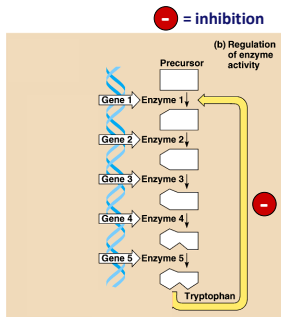


- Bacteria need to respond quickly to changes in their environment
 - if they have enough of a product, need to stop production
 - why?** waste of energy to produce more
 - how?** stop production of enzymes for synthesis
 - if they find new food/energy source, need to utilize it quickly
 - why?** metabolism, growth, reproduction
 - how?** start production of enzymes for digestion



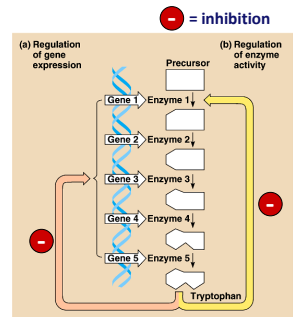
Remember Regulating Metabolism?

- Feedback inhibition
 - product acts as an **allosteric inhibitor** of 1st enzyme in tryptophan pathway
 - but this is wasteful production of enzymes*



Different way to Regulate Metabolism

- Gene regulation
 - instead of blocking enzyme function, block transcription of genes for all enzymes in tryptophan pathway
 - saves energy by not wasting it on unnecessary protein synthesis



Gene regulation in bacteria

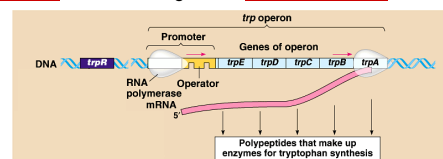


- Cells can vary the amount of specific enzymes by **regulating gene transcription**
 - turn **genes on** or turn **genes off**
 - turn genes OFF example**
 - if bacterium has enough tryptophan then it doesn't need to make enzymes used to **build** tryptophan
 - turn genes ON example**
 - if bacterium encounters new sugar (energy source), like lactose, then it needs to start making enzymes used to **digest** lactose



Bacteria group genes together

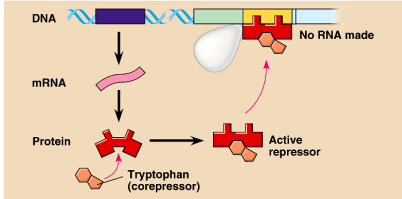
- Operon**
 - genes grouped together with related functions
 - example:** all enzymes in a metabolic pathway
 - promoter** = RNA polymerase binding site
 - single promoter controls transcription of all genes in operon
 - transcribed as **one unit** & a single mRNA is made
 - operator** = DNA binding site of **repressor protein**



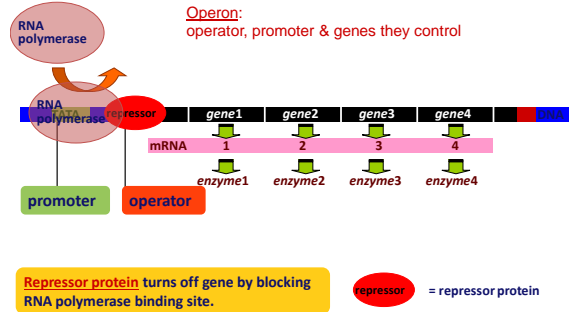
So how can these genes be turned off?

Repressor protein

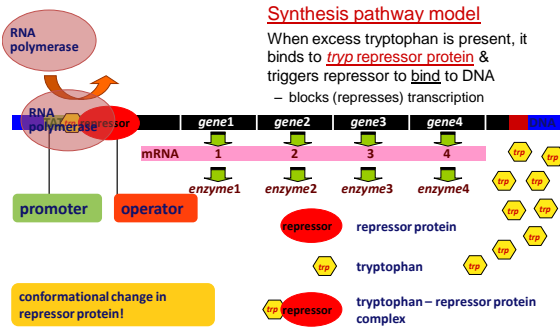
- binds to DNA at operator site
- blocking RNA polymerase
- blocks transcription



Operon Model

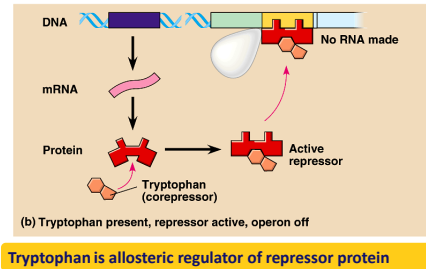


Repressible operon: tryptophan

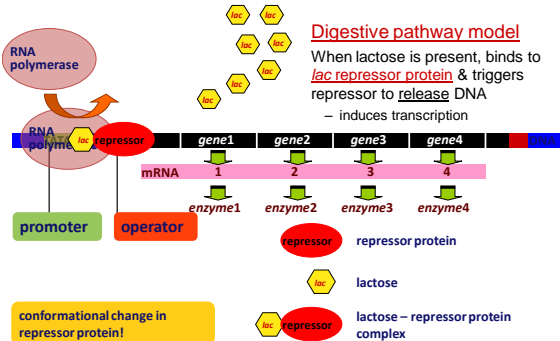


Tryptophan operon

What happens when tryptophan is present?
Don't need to make tryptophan-building enzymes

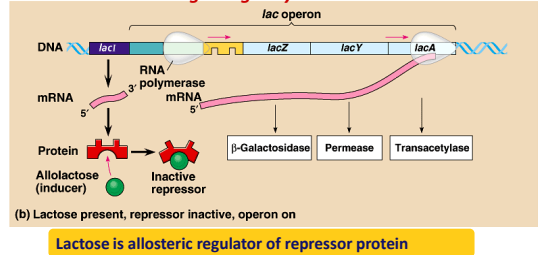


Inducible operon: lactose



Lactose operon

What happens when lactose is present?
Need to make lactose-digesting enzymes

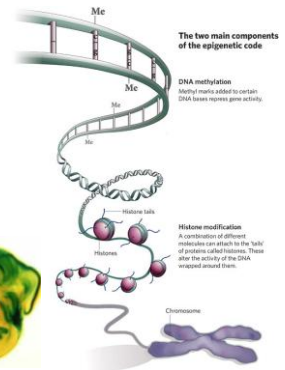
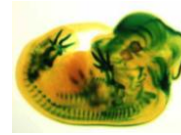


Operon summary



- **Repressible operon** (Trp Operon)
 - usually functions in **anabolic** pathways
 - **synthesizing** end products
 - when end product is present in excess, cell allocates resources to other uses
- **Inducible operon** (Lac Operon)
 - usually functions in **catabolic** pathways,
 - **digesting** nutrients to simpler molecules
 - produce enzymes only when nutrient is available
 - cell avoids making proteins that have nothing to do, cell allocates resources to other uses

Control of Eukaryotic Genes



The BIG Questions...

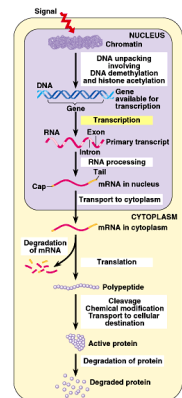


- How are genes turned on & off in eukaryotes?
- How do cells with the same genes differentiate to perform completely different, specialized functions?



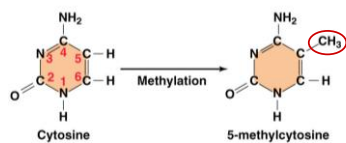
Points of control

- The control of gene expression can occur at any step in the pathway from gene to functional protein
 1. packing/unpacking DNA
 2. transcription
 3. mRNA processing
 4. mRNA transport
 5. translation
 6. protein processing
 7. protein degradation



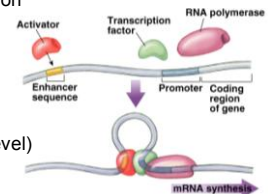
DNA methylation

- **Methylation of DNA** blocks transcription factors
 - no transcription
 - **genes turned off**
 - attachment of methyl groups (-CH₃) to cytosine
 - C = cytosine
 - nearly permanent inactivation of genes
 - ex. inactivated mammalian X chromosome = Barr body

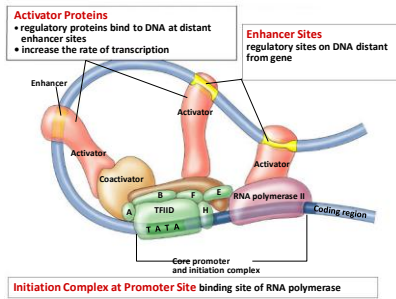


Transcription initiation

- Control regions on DNA
 - **promoter**
 - **nearby** control sequence on DNA
 - binding of RNA polymerase & **transcription factors**
 - “base” rate of transcription
 - **enhancer**
 - **distant** control sequences on DNA
 - binding of activator proteins
 - “enhanced” rate (high level) of transcription

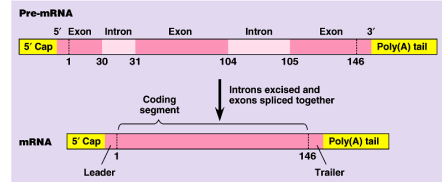


Transcription complex



Regulation of mRNA degradation

- 5' cap and Poly-A Tail
- Life span of mRNA determines amount of protein synthesis
 - mRNA can last from hours to weeks



RNA interference



- **Small interfering RNAs (siRNA)**
 - short segments of RNA (21-28 bases)
 - bind to mRNA
 - create sections of double-stranded mRNA
 - “death” tag for mRNA
 - triggers degradation of mRNA
 - cause **gene “silencing”**
 - post-transcriptional control
 - turns off gene = no protein produced

siRNA



1990s | 2006

RNA interference

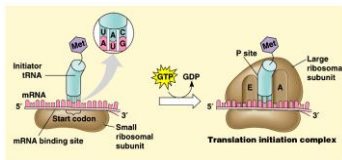
Andrew Fire Stanford
 Craig Mello U Mass

Nobel Prize

“for their discovery of RNA interference – gene silencing by double-stranded RNA”

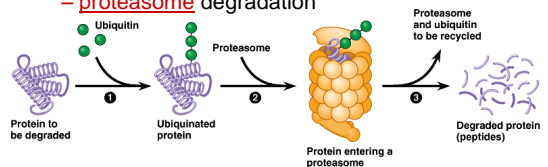
Control of translation

- **Block initiation of translation stage**
 - regulatory proteins attach to 5' end of mRNA
 - prevent attachment of ribosomal subunits & initiator tRNA
 - block translation of mRNA to protein



Protein processing & degradation

- Protein processing
 - folding, cleaving, adding sugar groups, targeting for transport
- Protein degradation
 - **ubiquitin tagging**
 - **proteasome** degradation



1980s | 2004

Ubiquitin

- “Death tag”
 - mark unwanted proteins with a label
 - 76 amino acid polypeptide, ubiquitin
 - labeled proteins are broken down rapidly in “waste disposers”
- proteasomes



Aaron Ciechanover
Israel



Avram Hershko
Israel

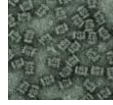


Irwin Rose
UC Riverside

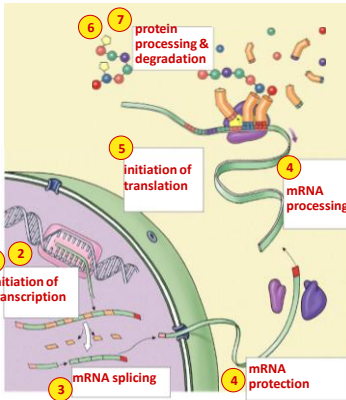


Proteasome

- Protein-degrading “machine”
 - cell’s waste disposer
 - breaks down any proteins into 7-9 amino acid fragments
 - cellular recycling



play Nobel animation



Gene Regulation

- 1 & 2. transcription**
 - DNA packing
 - transcription factors
- 3 & 4. post-transcription**
 - mRNA processing
 - splicing
 - 5' cap & poly-A tail
 - breakdown by siRNA
- 5. translation**
 - block start of translation
- 6 & 7. post-translation**
 - protein processing
 - protein degradation