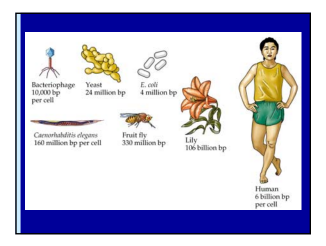
Lecture Series 8 The Eukaryotic Genome and Its Expression

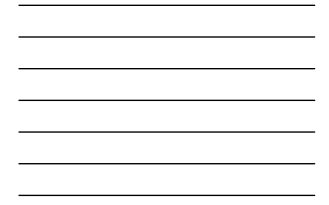
Reading Assignments

- Read Chapter 8
- Control of Gene Expression
- Skim Chapter 9
 How Genes and Genomes Evolve

A. The Eukaryotic Genome

 Although eukaryotes have more DNA in their genomes than prokaryotes, in some cases there is NO apparent relationship between genome size and organism complexity.







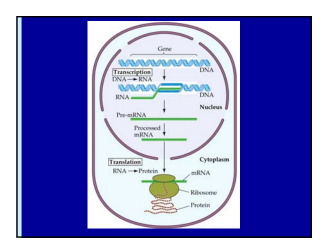
Amoeba dubia is the big winner at **670 Billion** base pairs per cell and an uncertain phylogeny!

CHARACTERISTIC	PROKARYOTES	EUKARYOTES
Genome size (base pairs)	104-107	108-1011
Repeated sequences	Few	Many
Noncoding DNA within coding sequences	Rare	Common
Transcription and translation separated in cell	No	Yes
DNA segregated within a nucleus	No	Yes
DNA bound to proteins	Some	Extensive
Promoter	Yes	Yes
Enhancer/silencer	Rare	Common
Capping and tailing of mRNA	No	Yes
RNA splicing required	Rare	Common
Number of chromosomes		
in genome	One	Many



A. The Eukaryotic Genome

- Unlike prokaryotic DNA, eukaryotic DNA is separated from the cytoplasm by being contained within a nucleus.
- The initial mRNA transcript of the DNA gets modified before it is exported to the cytoplasm.



A. The Eukaryotic Genome

• The genome of the single-celled budding yeast contains genes for the same metabolic machinery as bacteria, as well as genes for protein targeting in the cell.

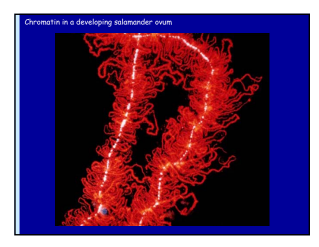
17.2 Comparison of the Ge	14.2 Comparison of the Genomes of E. coli and Yeast			
	E.COLI	S YEAST		
Genome length (base pairs)	4,640,000	12,068,000		
Number of proteins	4,300	6,200		
Proteins with roles in:				
Metabolism	650	650		
Energy production/storage	240	175		
Membrane transporters	280	250		
DNA replication/repair/ recombination	120	175		
Transcription	230	400		
Translation	180	350		
Protein targeting/secretion	35	430		
Cell structure	180	250		

A. The Eukaryotic Genome

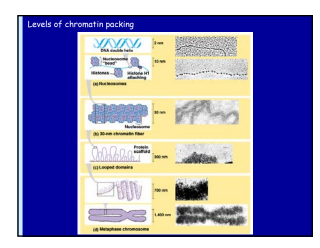
- The genome of the multicellular roundworm *Caenorhabditis elegans* contains genes required for intercellular interactions.
- The genome of the fruit fly has fewer genes than that of the roundworm. Many of its sequences are homologs of sequences on roundworm and mammalian genes.

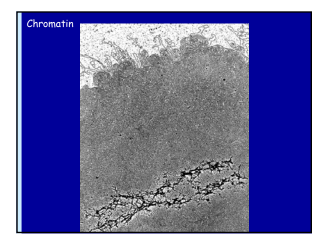
FUNCTION	PROTEIN/DOMAIN	GENES	
Transcription control	Zinc finger; homeobox	540	
RNA processing	RNA binding domains	100	
Nerve impulse transmission	Gated ion channels	80	
Tissue formation	Collagens	170	
Cell interactions	Extracellular domains; glycotransferases	330	
Cell-cell signaling	G protein-linked receptors; protein	1,290	
	kinases; protein phosphatases		

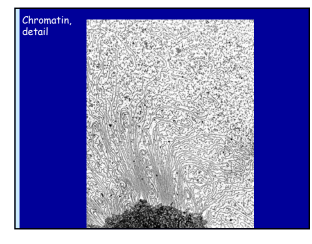












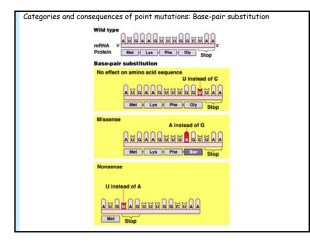


B. Mutations: Heritable Changes in Genes

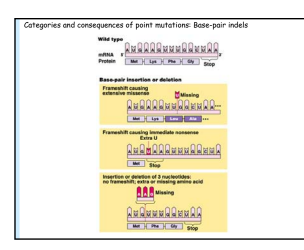
- Mutations in DNA are often expressed as abnormal proteins. However, the result may not be easily observable phenotypic changes.
- Raw materials for evolution to operate.
- Some mutations appear only under certain conditions, such as exposure to a certain environmental agent or condition.

B. Mutations: Heritable Changes in Genes

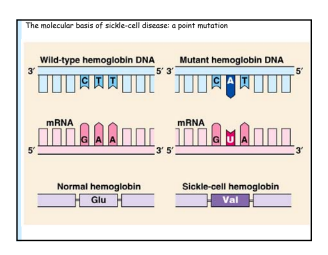
 Point mutations (silent, missense, nonsense, or frame-shift) result from alterations in single base pairs of DNA.







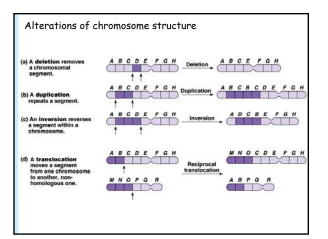






B. Mutations: Heritable Changes in Genes

 Chromosomal mutations (deletions, duplications, inversions, or translocations) involve large regions of a chromosome.

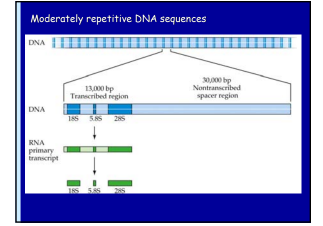


C. Repetitive Sequences

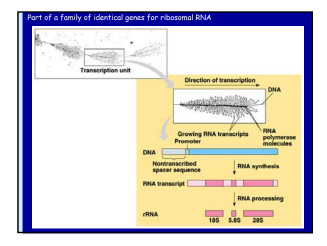
- Highly repetitive DNA is present in up to millions of copies of short sequences. It is not transcribed. Its role is unknown.
- Rem: Some moderately repetitive DNA sequences, such as telomeric DNA is found at the ends of chromosomes.

C. Repetitive Sequences

- Some moderately repetitive DNA sequences, such as those coding for ribosomal RNA's, are transcribed.
- Up to three rRNAs result, two go to the large subunit and one goes to the small subunit.



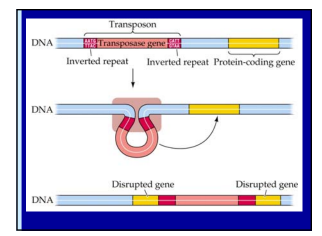






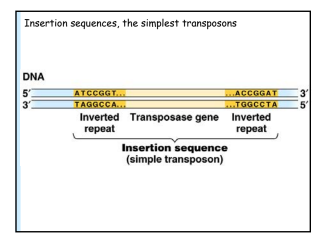
C. Repetitive Sequences

- Some moderately repetitive DNA sequences are transposable, or able to move about the genome. These are known as Transposons.
- Transposons can jump from place to place on the chromosome by actually moving or by making a new copy, inserted at a new location.

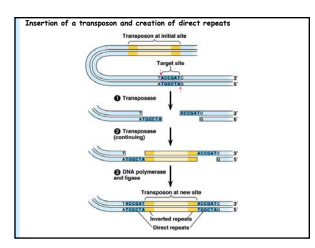




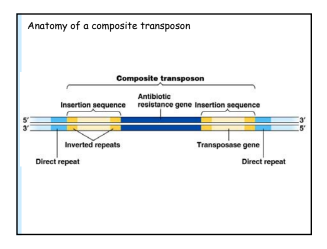




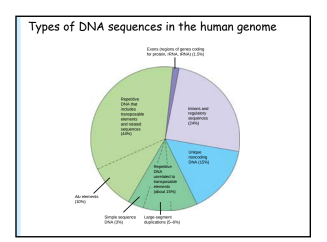








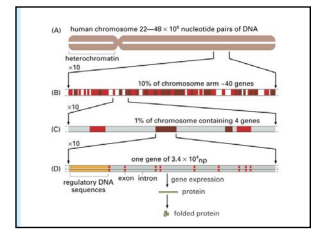




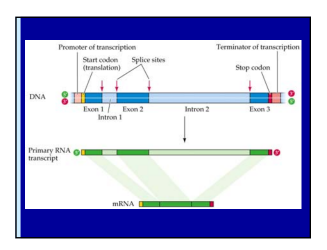


D. The Structures of Protein-Coding Genes

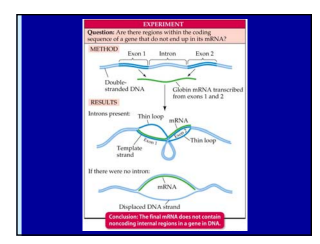
- A typical protein-coding gene has noncoding internal sequences (introns) as well as flanking sequences that are involved in the machinery of transcription and translation in addition to its exons or coding regions.
- These are usually single copy genes.







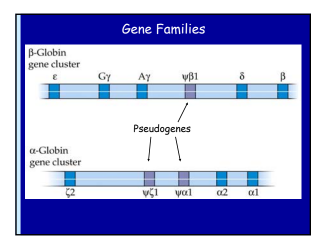




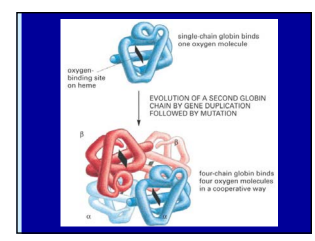


D. The Structures of Protein-Coding Genes

- Some eukaryotic genes form families of related genes that have similar sequences and code for similar proteins. These related proteins may be made at different times and in different tissues.
- Some sequences in gene families are pseudogenes, which code for nonfunctional mRNA's or proteins.

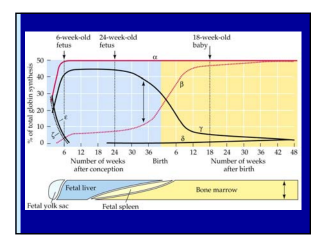




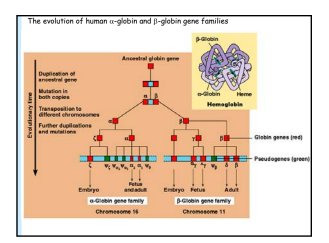


D. The Structures of Protein-Coding Genes

• Differential expression of different genes in the β -globin family ensures important physiological changes during human development.



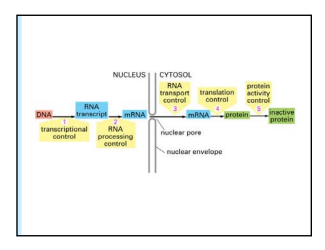






E. Transcriptional Control

 Eukaryotic gene expression can be controlled at the transcriptional, posttranscriptional, translational, and posttranslational levels.



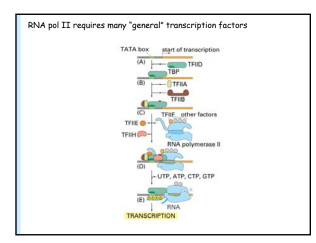


E. Transcriptional Control

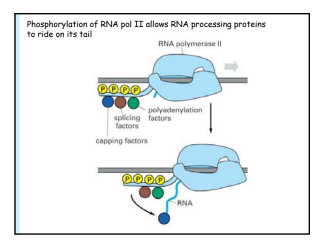
• The major method of control of eukaryotic gene expression is selective transcription, which results from specific proteins binding to regulatory regions on DNA.

E. Transcriptional Control

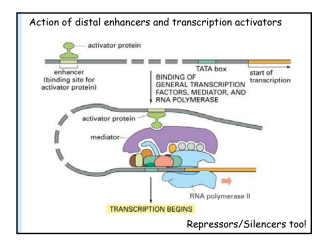
- A series of "general" transcription factors must bind to the promoter before RNA polymerase can bind.
- Whether RNA polymerase will initiate transcription also depends on the binding of regulatory proteins, activator proteins, and repressor proteins.







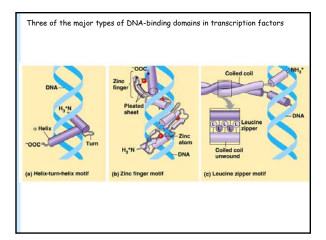






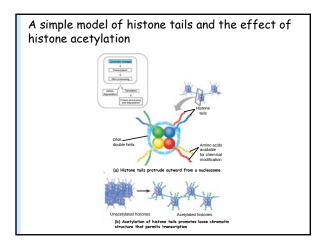
E. Transcriptional Control

 The DNA-binding domains of most DNAbinding proteins have one of four structural motifs: helix-turn-helix, zinc finger, leucine zipper, or homeodomain.

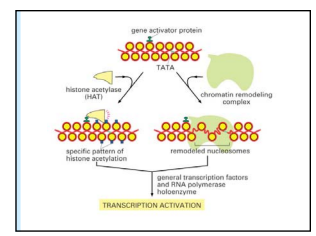


E. Transcriptional Control

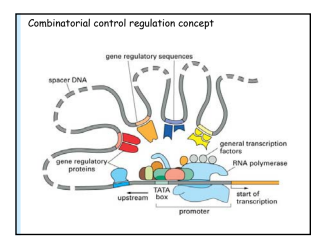
• Acetylation of histone tails promotes loose chromatin structure that permits transcription to more readily occur.







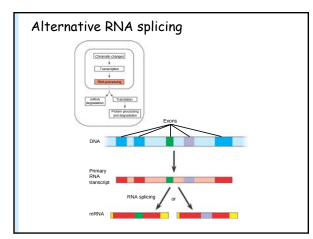






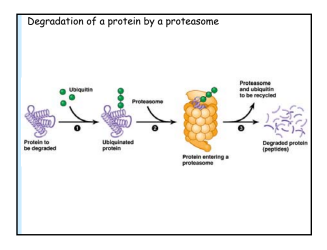
F. Posttranscriptional Control

- Because eukaryotic genes have several exons, alterative mRNAs can be generated from the same RNA transcript.
- This alternate splicing can be used to produce different proteins.
- The stability of mRNA in the cytoplasm can be regulated by the binding of proteins.

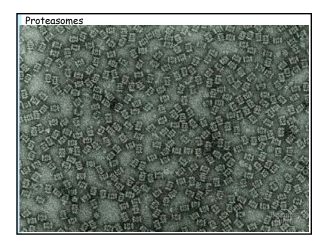


F. Posttranslational Control

• Proteasomes degrade proteins targeted for breakdown.

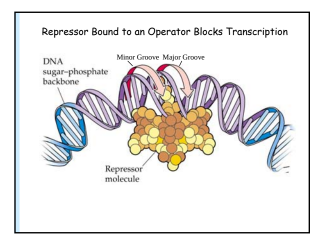






G. Regulation of Gene Expression in Prokaryotes

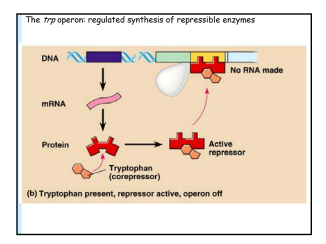
- An operon consists of a promoter, an operator, and structural genes. Promoters and operators do not code for proteins, but serve as binding sites for regulatory proteins.
- When a repressor protein binds to the operator, transcription of the structural genes is inhibited.



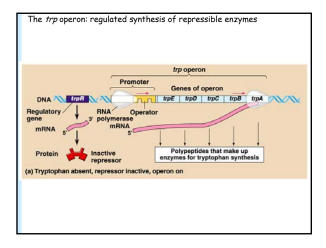


G. Regulation of Gene Expression in Prokaryotes

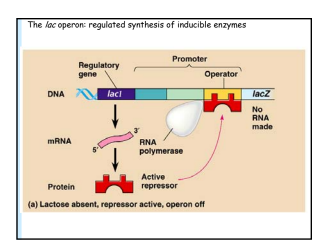
- The expression of prokaryotic genes is regulated by: inducible operator-repressor systems, repressible operator-repressor systems (e.g., both negative control), and systems that increase the efficiency of a promoter (e.g., positive control).
- Repressor proteins are coded by constitutive regulatory genes.



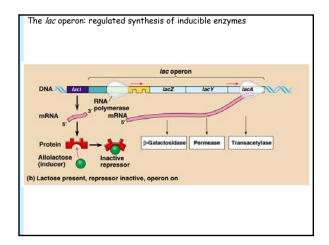








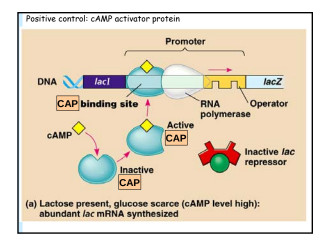




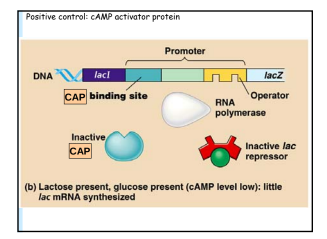


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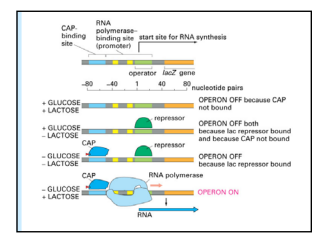
- The efficiency of RNA polymerase can be increased by regulation of the level of cyclic AMP, which binds to CAP (cAMP activator protein).
- The CAP-cAMP complex then binds to a site near the promoter of a target gene, enhancing the binding of RNA polymerase and hence transcription.













13.2	The Relat	tionships Between	Positive and	d Negative Control in ti	he lac Operon		
GLUCOSE	cAMP LEVELS	RNA POLYMERASE BINDING TO PROMOTER	LACTOSE	LAC REPRESSOR	TRANSCRIPTION OF LAC GENES?	LACTOSE USED BY CELLS	
Present	Low	Absent	Absent	Active and bound to operator	No	No	
Present	Low	Absent	Present	Inactive and not bound to operator	No	No	
Absent	High	Present	Present	Inactive and not bound to operator	Yes	Yes	
Absent	High	Absent	Absent	Active and bound to operator	No	No	



H. Comparison of Control Features in Bacteria & Eucarya

- Bacteria have multiple genes under single control: operons
- Eucarya have multiple RNA polymerases
- Simple vs. Complex Transcription Factors
- Local vs. Distal Control: Enhancers/Silencers
- Eucarya must contend with Chromatin