

Regulation of gene expression in eukaryotes

Major principle: Activation of gene activity

Positive Control of Gene expression

General Chromatin structure

Wide domain regulators

Gene-specific Regulators

Coregulators

Modification of regulators



Translation - Eukaryotes

Start Codon

mRNA 5'-CAP.....AUG

Influences:

Surrounding of AUG!!!

Kozak Consensus

......CC^A/_GCCAUGG..... mammalian

..... $A/_TA^A/_CA^A/_CAAUGTC^T/_C$ Yeast

...... gccgcc(A/G)ccAUGG Wikipedia



Combinatorial Principle II Cell type/Tissue В Developmental stage all B A B



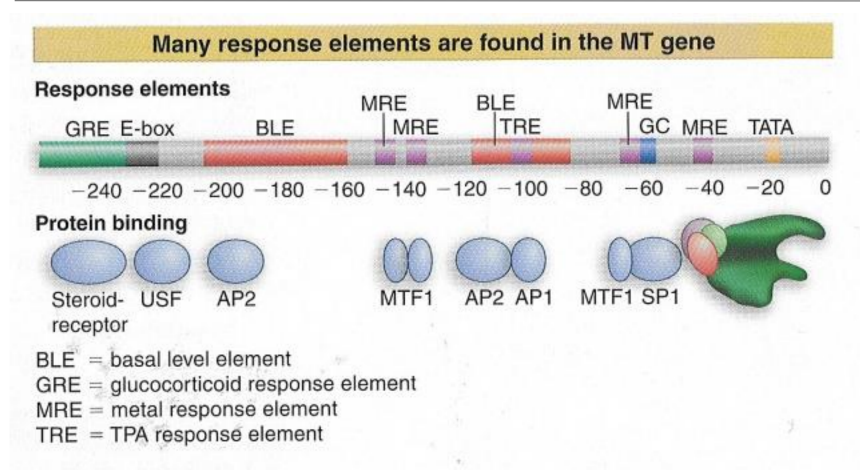


Figure 25.7 The regulatory region of a human metallothionein gene contains regulator elements in both its promoter and enhancer. The promoter has elements for metal induction; an enhancer has an element for response to glucocorticoid. Promoter elements are shown above the map, and proteins that bind them are indicated below.



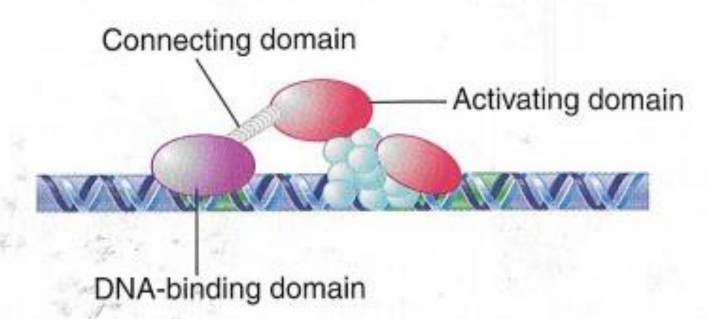


FIGURE 28.6 DNA-binding and activating functions in a transcription factor may comprise independent domains of the protein.



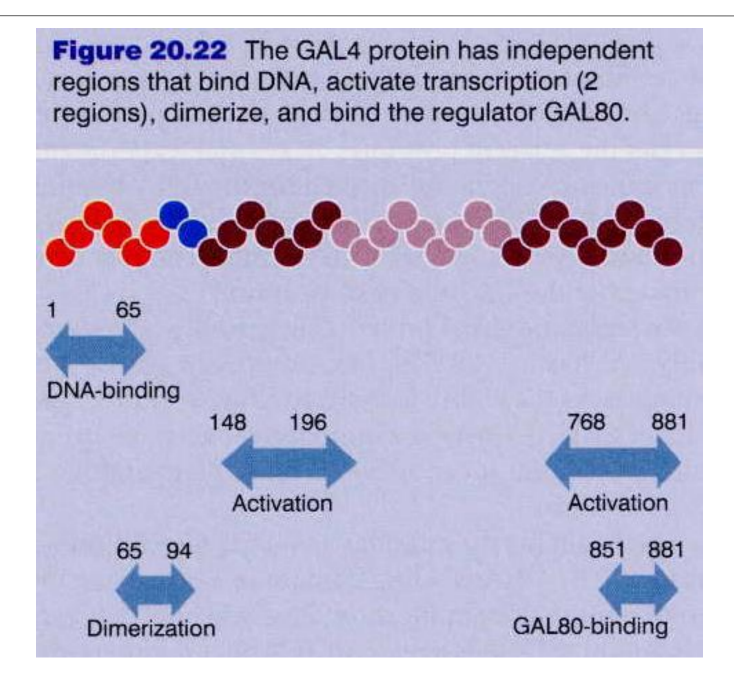




Figure 20.23 The ability of GAL4 to activate transcription is independent of its specificity for binding DNA. When the GAL4 DNA-binding domain is replaced by the LexA DNA-binding domain, the hybrid protein can activate transcription when a LexA operator is placed near a promoter. UASG Promoter LexA target Promoter **GAL4** activator Binding & transcription No binding **GAL4 DNA-binding GAL4** activator Binding & transcription No binding LexA DNA-binding



Figure 20.24 The activating domain of the tat protein of HIV can stimulate initiation if it is tethered in the vicinity by binding to the RNA product of a previous round of transcription. Activation is independent of the means of tethering, as shown by the substitution of a DNA-binding domain for the RNA-binding domain. Usually tat binds to the tar sequence in the RNA product RNA-binding domain Activation domain Transcript Initiation complex Transcribing polymerase A tat activation domain linked to a DNA-binding sequence works just as well Activation domain **DNA-binding domain**



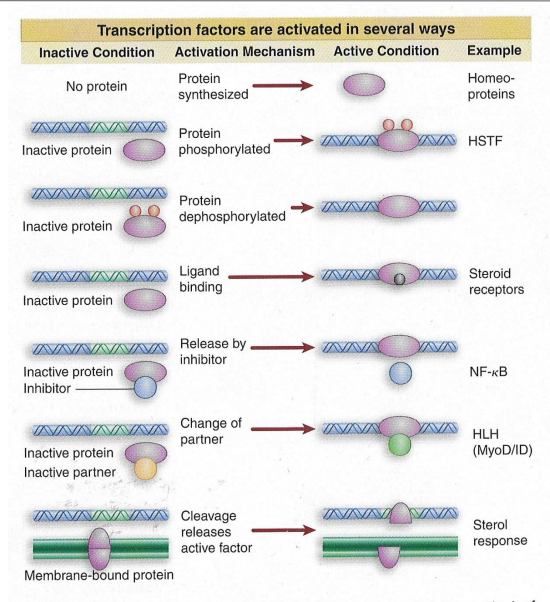


Figure 25.8 The activity of a regulatory transcription factor may be controlled by synthesis of protein, covalent modification of protein, ligand binding, or binding of inhibitors that sequester the protein or affect its ability to bind to DNA.

Taken from: B. Lewin, Essential Genes, Pearson Ed. International



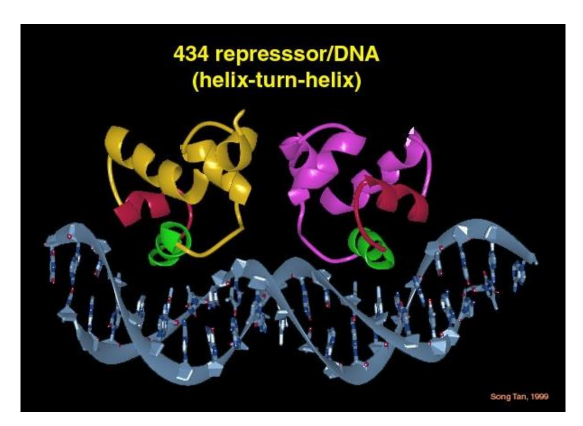
DNA Binding Proteins - motifs



Helix –Turn – Helix Proteins

The Helix-Turn-Helix motif consists of two a helices and a short extended amino acid chain between them. The more carboxyl-terminal helix can fit into the major groove of DNA. This motif is found in hundreds of DNA-binding proteins, including λ - repressor, tryptophan repressor, catabolite activator protein (CAP), octamer transcription factor 1 (Oct-1) and heat shock factor (HSF).

Source: http://www.web-books.com/MoBio/Free/Ch4F4.htm



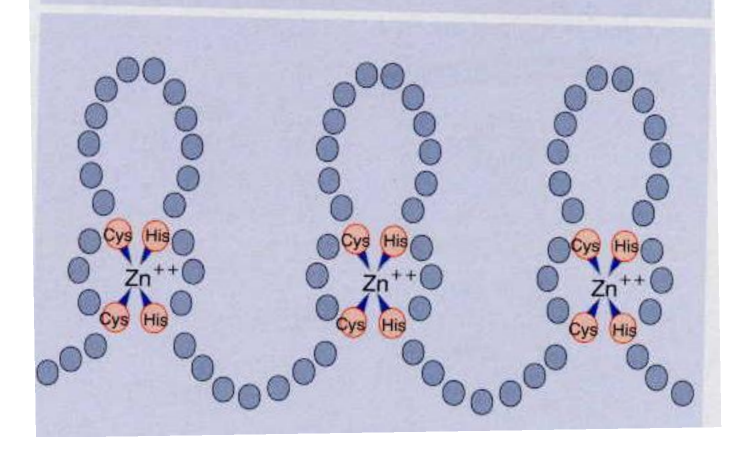
Source: http://www.bmb.psu.edu/faculty/tan/lab/gallery/434reprdna.jpg

See also: http://www.proteopedia.org/wiki/index.php/Helix-turn-helix_motif



Zink-Finger Proteins

Figure 21.3 Transcription factor SP1 has a series of three zinc fingers, each with a characteristic pattern of cysteine and histidine residues that constitute the zinc-binding site.





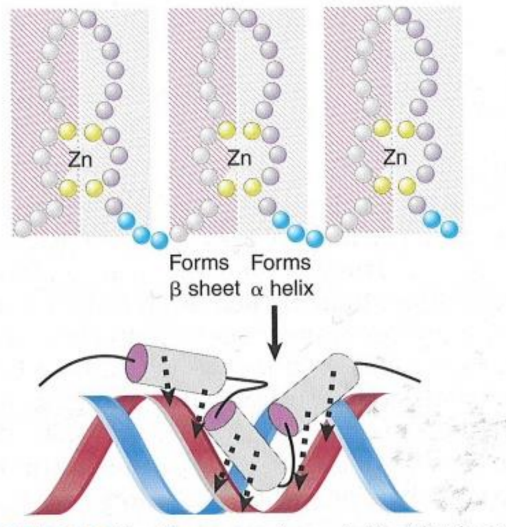


FIGURE 28.12 Zinc fingers may form α helices that insert into the major groove, which is associated with β sheets on the other side.



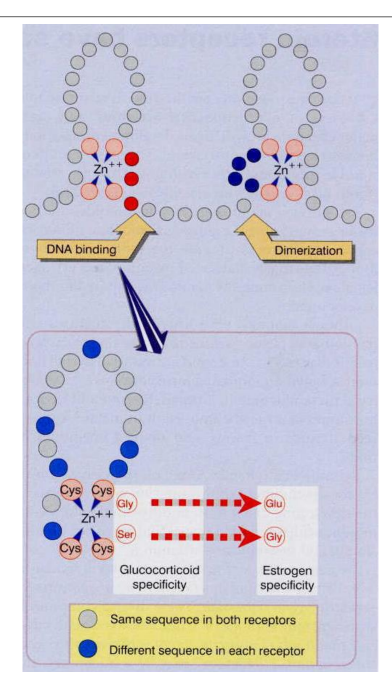


Figure 21.5 The first finger of a steroid receptor controls specificity of DNA-binding (positions shown in red); the second finger controls specificity of dimerization (positions shown in blue). The expanded view of the first finger shows that discrimination between GRE and ERE target sequences rests on two amino acids at the base.



Ligand-gated receptors share structural features

DNA binding and transcriptional activation (identity varies from 94–42%)

N-terminal regions have <15% identities (needed to activate transcription) Hormone-binding regions and dimerization (identity varies from 57–15%)

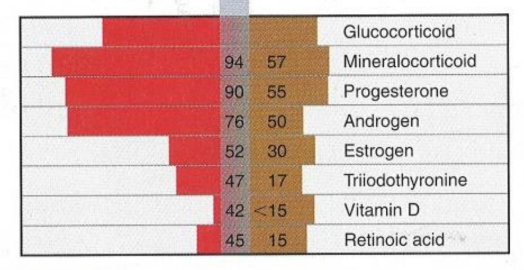


Figure 25.12 Receptors for many steroid and thyroid hormones have a similar organization, with an individual N-terminal region, conserved DNA-binding region, and a C-terminal hormone-binding region. Identities are relative to GR.



Repression prevails in absence of ligand

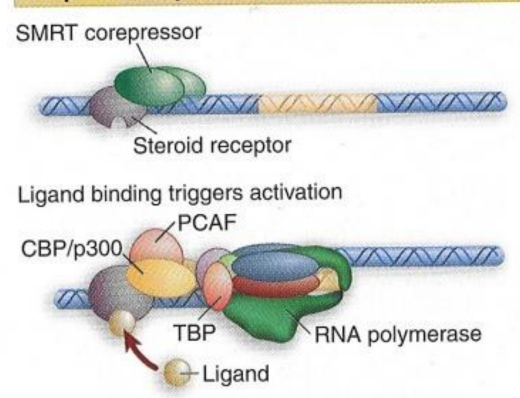


Figure 25.17 The steroid receptors TR and RAR bind the SMRT corepressor in the absence of ligand. The promoter is not expressed. When SMRT is displaced by binding of ligand, the receptor binds a coactivator complex. This leads to activation of transcription by the basal apparatus.



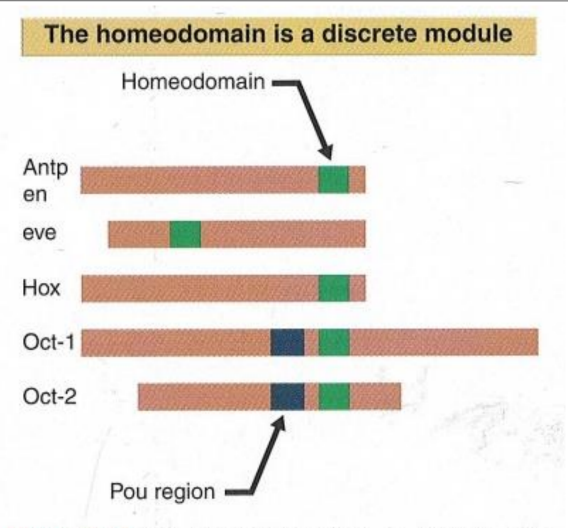


Figure 25.18 The homeodomain may be the sole DNAbinding motif in a transcriptional regulator or may be combined with other motifs. It represents a discrete (60 residue) part of the protein.



The homeodomain is a module of 60 amino acids N-terminal arm 10 Helix 1 20 1 Glu Lys Arg Pro Arg Thr Ala Phe Ser Ser Glu Gln Leu Ala Arg Leu Lys Arg Glu Phe Asn Glu En Arg Lys Arg Gly Arg Gln Thr Tyr Thr Arg Tyr Gln Thr Leu Glu Leu Glu Lys Glu Phe His Phe Antp Arg Arg Lys Lys Arg Thr Ser Ile Glu Thr Asn Val Arg Phe Ala Leu Glu Lys Ser Phe Leu Ala Oct2 30 Helix 2 40 En Asn Arg Tyr Leu Thr Glu Arg Arg Arg Glu Glu Leu Ser Ser Glu Leu Gly Leu Asn Arg Tyr Leu Thr Arg Arg Arg Ile Glu Ile Ala His Ala Leu Cys Leu Antp Asn Glu Lys Pro Thr Ser Glu Glu Ile Leu Leu Ile Ala Glu Gln Leu His Met Oct2 Helix 3 41 50 60 Asn Glu Ala Gln Ite Lys Ile Trp Phe Gln Asn Lys Arg Ala Lys Ile Lys Lys Ser Asn En Thr Glu Arg Gln Ile Lys Ile Trp Phe Gln Asn Arg Arg Met Lys Trp Lys Lys Glu Asn Antp Glu Lys Glu Val Ile Arg Val Trp Phe Cys Asn Arg Arg Gln Lys Glu Lys Arg Ile Asn Oct2

Figure 25.19 The homeodomain of the *Antennapedia* gene represents the major group of genes containing homeoboxes in *Drosophila*; *engrailed* (*en*) represents another type of homeotic gene; and the mammalian factor Oct2 represents a distantly related group of transcription factors. The homeodomain is conventionally numbered from 1 to 60. It starts with the N-terminal arm, and the three helical regions occupy residues 10–22, 28–38, and 42–58. Amino acids in red are conserved in all three examples.



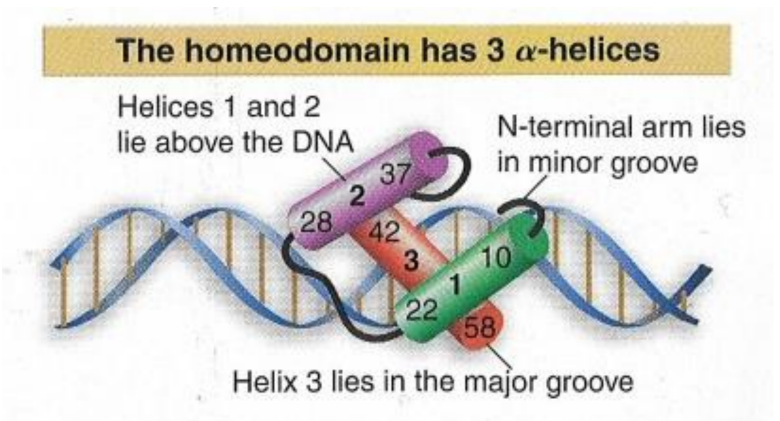


Figure 25.20 Helix 3 of the homeodomain binds in the major groove of DNA, with helices 1 and 2 lying outside the double helix. Helix 3 contacts both the phosphate backbone and specific bases. The N-terminal arm lies in the minor groove, and makes additional contacts.

ld



HLH proteins have two helical regions

Ala Asp Arg Arg Lys Ala Ala Thr Met Arg Gln Arg Arg Arg MyoD 6 conserved residues Arg Leu Pro Ala Leu Leu Asp Gln Glu Glu Val Asn Val Leu ld are absent from Id Helix 1 Leu Ser Lys Val Asn Gln Ala Phe Gln Thr Leu Lys Arg Cys Thr MyoD Conserved residues are Leu Tyr Asp Met Asn Gly Cys Tyr Ser Arg Leu Lys Gln Leu Val

Lys Val Gln Ile Leu Arg Asn Ala Ile Arg Tyr Ile Gln Gly Leu Glu MyoD Helix 2 Lys Val Gln Ile Leu Glu His Val Ile Asp Tyr Ile Arg Asp Leu Glu ld

Figure 25.21 All HLH proteins have regions corresponding to helix 1 and helix 2, separated by a loop of 10–24 residues. Basic HLH proteins have a region with conserved positive charges immediately adjacent to helix 1.

Basic region

found in both MyoD and Id



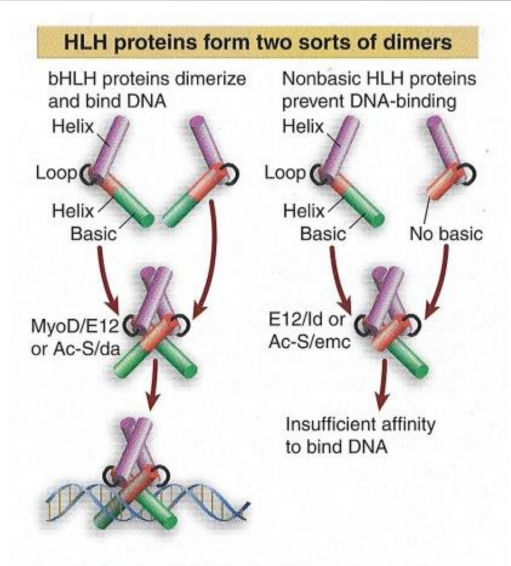


Figure 25.22 An HLH dimer in which both subunits are of the bHLH type can bind DNA, but a dimer in which one subunit lacks the basic region cannot bind DNA.



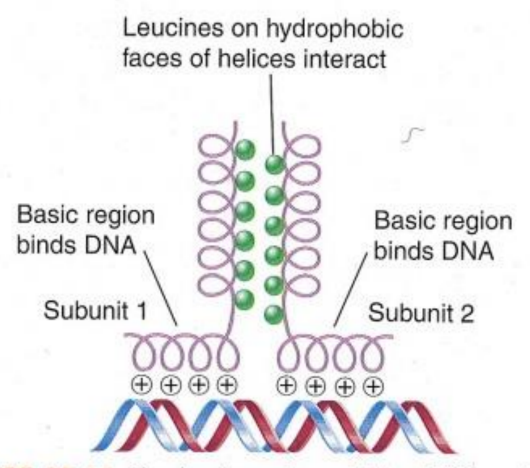


FIGURE 28.16 The basic regions of the bZIP motif are held together by the dimerization at the adjacent zipper region when the hydrophobic faces of two leucine zippers interact in parallel orientation.



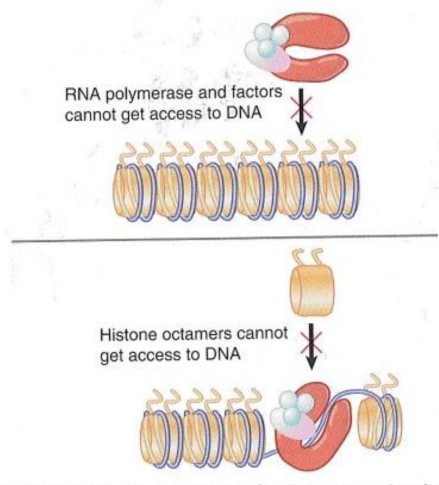


FIGURE 28.17 If nucleosomes form at a promoter, transcription factors (and RNA polymerase) cannot bind. If transcription factors (and RNA polymerase) bind to the promoter to establish a stable complex for initiation, histones are excluded.



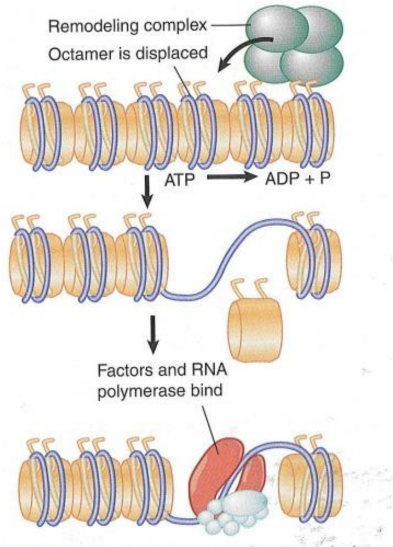


FIGURE 28.18 The dynamic model for transcription of chromatin relies upon factors that can use energy provided by hydrolysis of ATP to displace nucleosomes from specific DNA sequences.



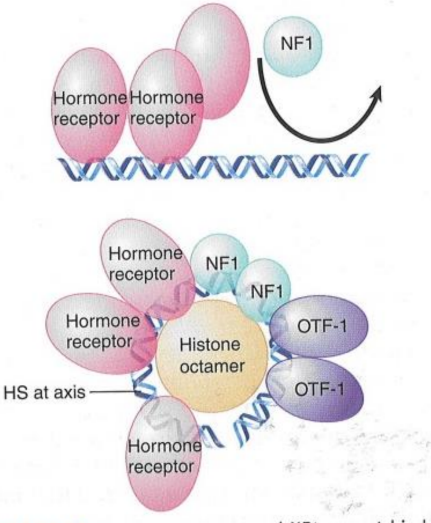


FIGURE 28.23 Hormone receptor and NF1 cannot bind simultaneously to the MMTV promoter in the form of linear DNA, but can bind when the DNA is presented on a nucleosomal surface.

Taken from: J.E. Krebs, E.S. Goldstein, S.T. Kilpatrick; "Lewin's Genes XI"; Jones&Bartlett Learning



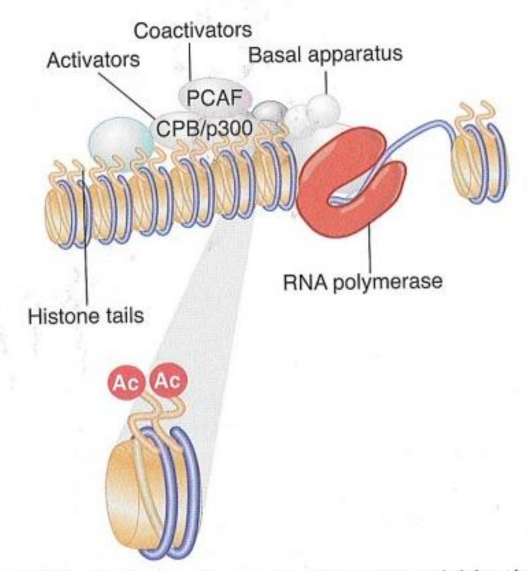


FIGURE 28.24 Coactivators may have HAT activities that acetylate the tails of nucleosomal histones.



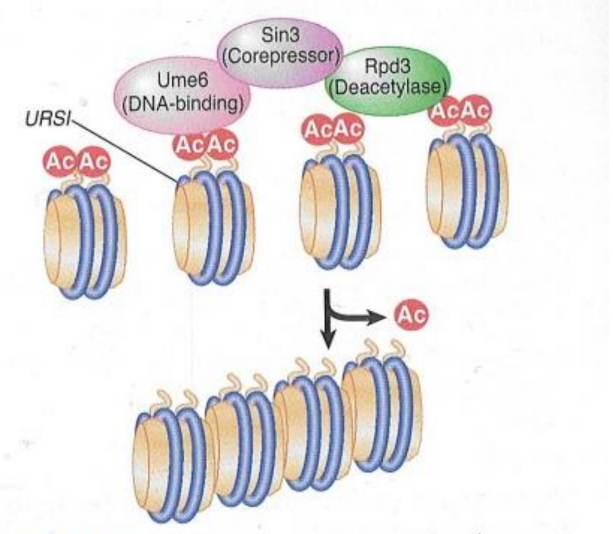


FIGURE 28.26 A repressor complex contains three components: a DNA-binding subunit, a corepressor, and a histone deacetylase.



Figure 21.6 Several types of hydrophobic small molecules activate transcription factors. Corticoids and steroid sex hormones are synthesized from cholesterol, vitamin D is a steroid, thyroid hormones are synthesized from tyrosine, and retinoic acid is synthesized from isoprene (in fish liver).

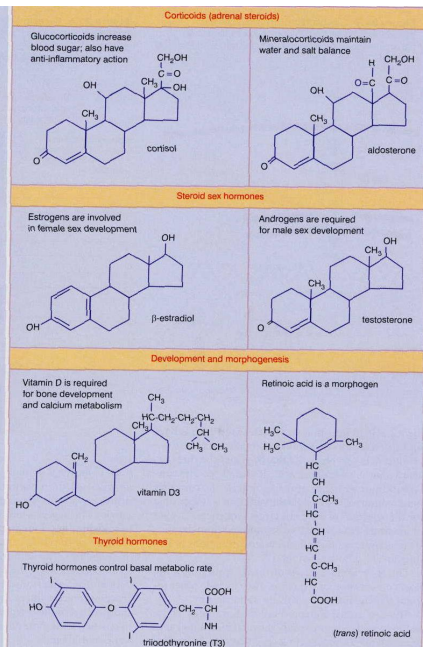


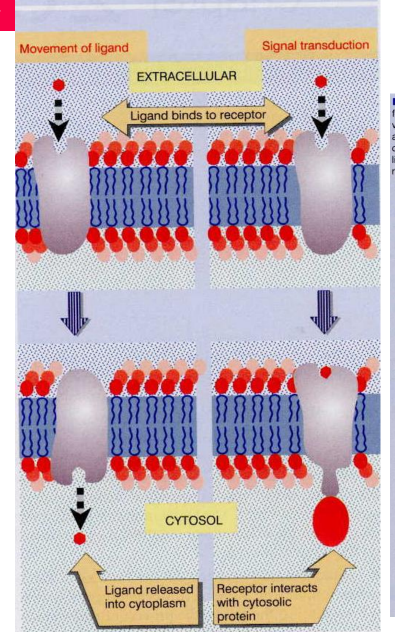


Figure 21.7 Glucocorticoids regulate gene transcription by causing their receptor to bind to an enhancer whose action is needed for promoter function. Steroid Receptor Cytoplasm Nucleus GRE/Enhancer Promoter Transcription Activation



29.11.16

Figure 26.1 Overview: information may be transmitted from the exterior to the interior of the cell by movement of a ligand or by signal transduction.





Signal transduction

Figure 26.2 Three means for transferring material of various sizes into the cell are provided by ion channels, receptor-mediated ligand transport, and receptor internalization.

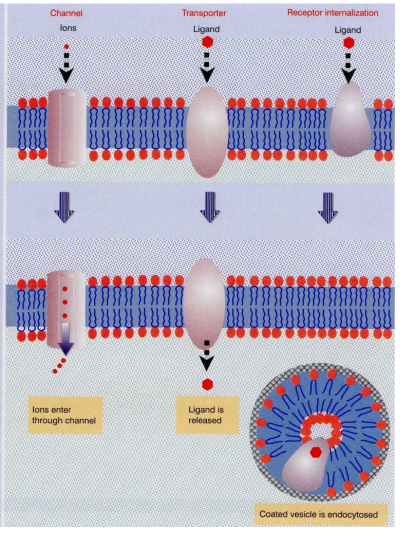




Figure 26.3 A signal may be transduced by activating the kinase activity of the cytoplasmic domain of a transmembrane receptor or by dissociating a G protein into subunits that act on target proteins on the membrane.

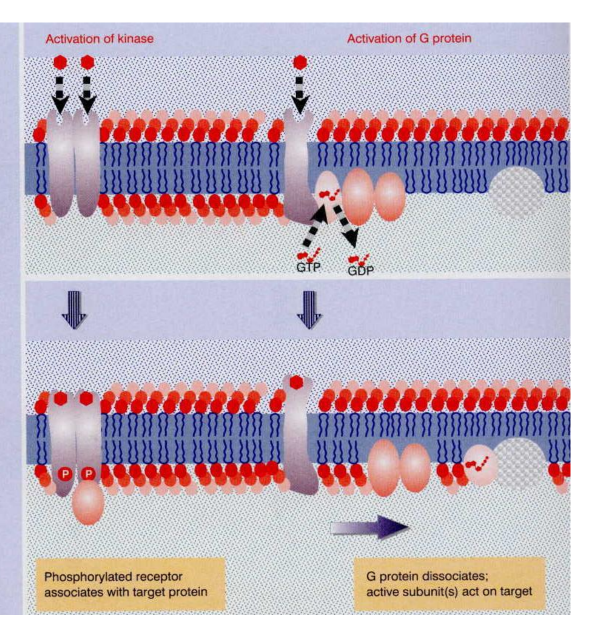




Figure 26.13 The principle underlying signal transduction by a tyrosine kinase receptor is that ligand binding to the extracellular domain triggers dimerization; this causes a conformational change in the cytoplasmic domain that activates the tyrosine kinase catalytic activity.

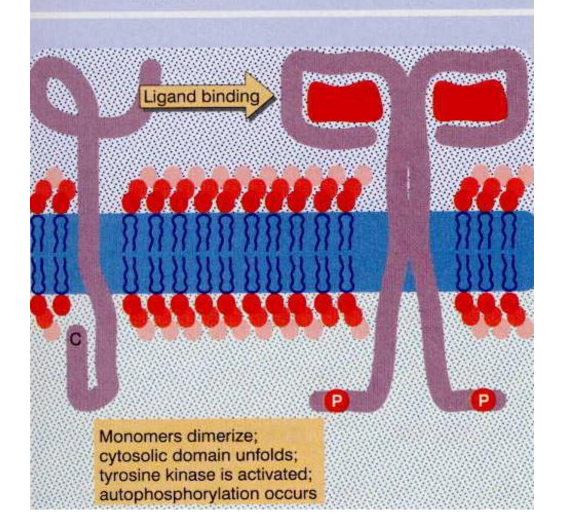




Figure 26.12 Effectors for receptor tyrosine kinases include phospholipases and kinases that act on lipids to generate second messengers.

Effector	Substrate	Products
PLC (phospolipase C) (3 families, PLC α , β , γ)	PIP2 (phosphatidylinositol 4,5-diphosphate)	DAG (diacylglycerol) + IP3 (inositol 1,4,5-triphosphate) DAG activates protein kinase C IP3 mobilizes Ca ²⁺
PLA2 (phospholipase A2)	Phospholipids (phosphatidylcholine, phosphatidylethanolamine, phosphatidylinositol)	Arachidonic acid Converted to prostaglandins & leukotrienes
PI3 kinase (phosphatidylinositol-3 kinase)	Phosphatidyl inositol	PI3 (phosphatidyl inositol-3 phosphate)
PI4 kinase (phosphatidylinositol-4 kinase)	Phosphatidyl inositol	PI4 (phosphatidyl inositol-4 phosphate) Converted to PIP2 (phosphatidyl diphosphate)

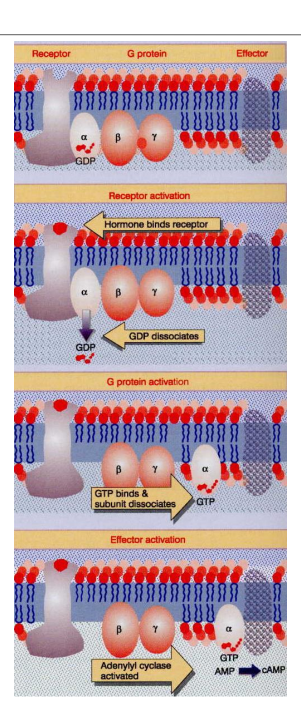


Figure 26.11 When a receptor is activated by hormone binding, it causes GTP to replace GDP on a $G\alpha$ subunit. The $G\alpha$ subunit dissociates from the $\beta\gamma$ dimer, and activates an effector such as adenylate cyclase.



Determination of Gene Function by DNA Rearrangements



Figure 24.17 Immunoglobulin type and function is determined by the heavy chain. J is a 'joining protein' in IgM; all other Ig types exist as tetramers.

Туре	IgM	IgD	lgG	IgA	IgE
Heavy chain	μ	δ	γ	α	ε
Structure	$(\mu_2 L_2)_5 J$	$\delta_2 L_2$	γ ₂ L ₂	$(\alpha_2 L_2)_2 J$	$\epsilon_2 L_2$
Proportion	5%	1%	80%	14%	<1%
Effector function	Activates	Development	Activates	Found in	Allergic
	complement	of tolerance (?)	complement	secretions	response



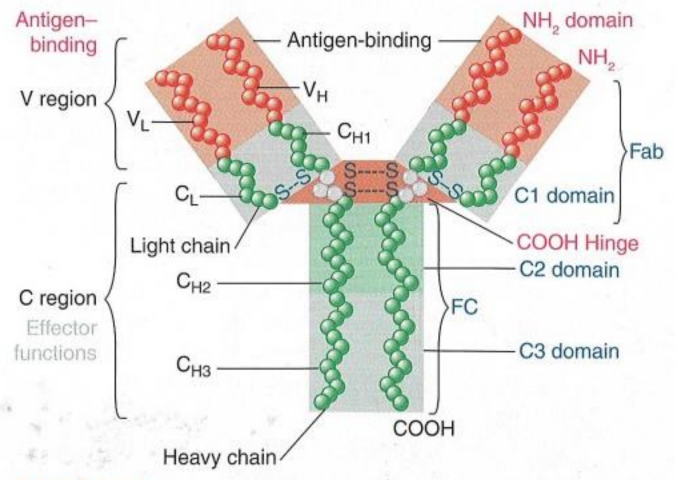


FIGURE 18.7 An antibody (immunoglobulin, or Ig) molecule is a heterodimer consisting of two identical heavy chains and two identical light chains. Schematized here is an IgG1, which comprises an N-terminal variable (V) region and a C-terminal constant (C) region.



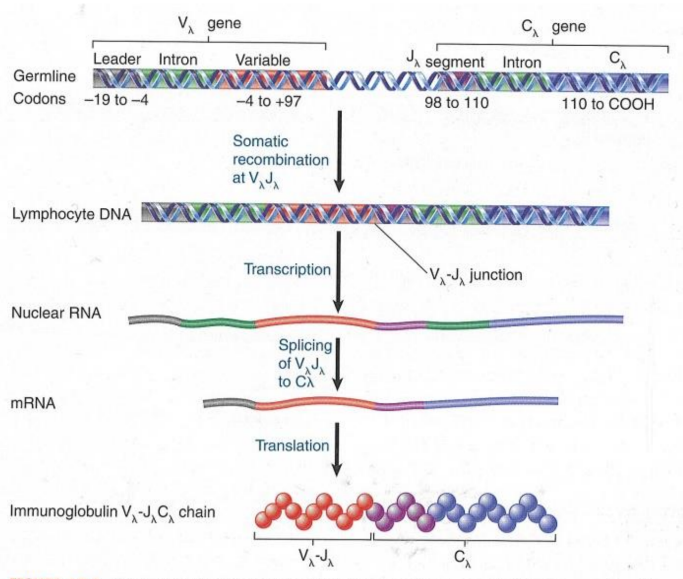


FIGURE 18.8 The C_{λ} gene segment is preceded by a J_{λ} segment, so that V_{λ} - J_{λ} recombination generates a productive V_{λ} - $J_{\lambda}C_{\lambda}$.

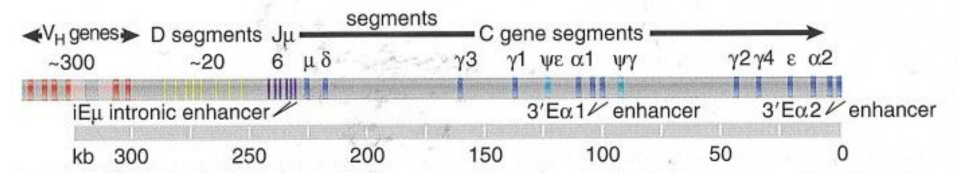


FIGURE 18.13 A single gene cluster in humans contains all the information for the IgH chain. Depicted is a schematic map of the human IgH chain locus.



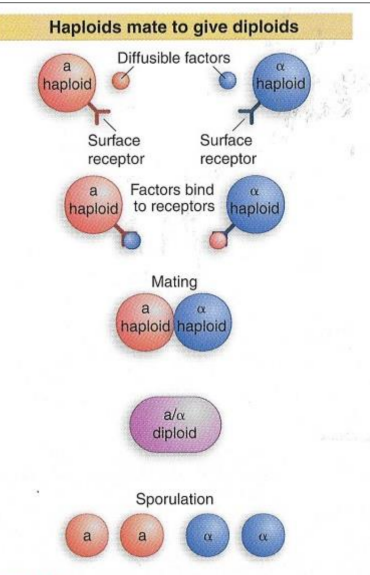


Figure 19.21 The yeast life cycle proceeds through mating of MATa and $MAT\alpha$ haploids to give heterozygous diploids that sporulate to generate haploid spores.

Regulation of expression by DNA rearrangements

Yeast mating type switching

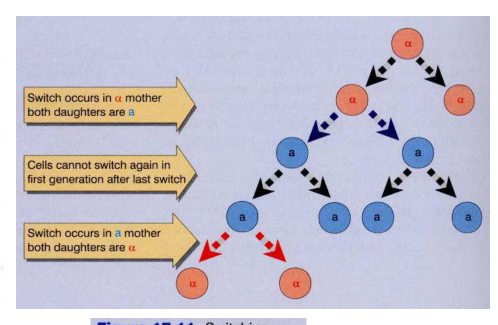
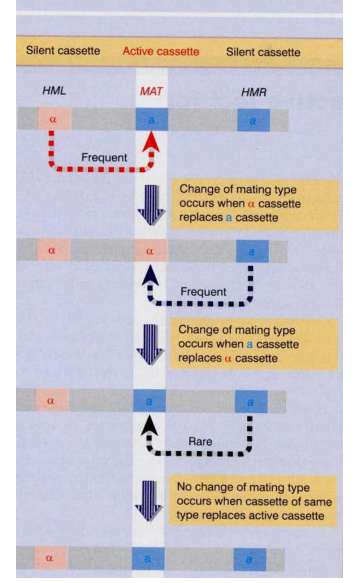


Figure 17.11 Switching occurs only in mother cells; both daughter cells have the new mating type. A daughter cell must pass through an entire cycle before it becomes a mother cell that is able to switch again.



Figure 17.5 Changes of mating type occur when silent cassettes replace active cassettes of opposite genotype; when transpositions occur between cassettes of the same type, the mating type remains unaltered.



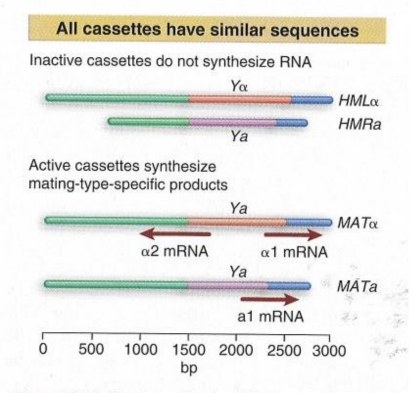


Figure 19.23 Silent cassettes have the same sequences as the corresponding active cassettes, except for the absence of the extreme flanking sequences in *HMRa*. Only the *Y* region changes between a and α types.



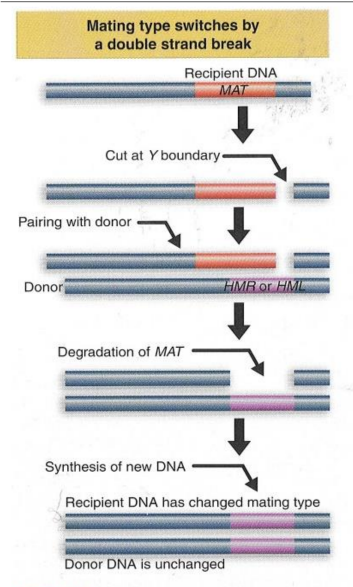


Figure 19.25 Cassette substitution is initiated by a double-strand break in the recipient (*MAT*) locus, and may involve pairing on either side of the *Y* region with the donor (*HMR* or *HML*) locus.

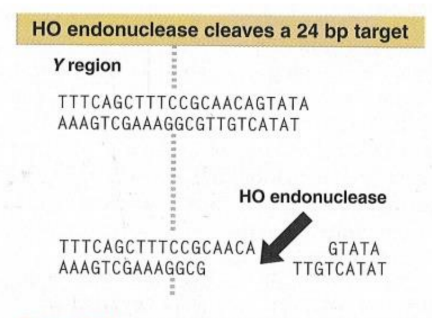


Figure 19.24 HO endonuclease cleaves *MAT* just to the right of the *Y* region, generating sticky ends with a 4-base overhang.



Figure 17.13 Overview: a trypanosome passes through several morphological forms when its life cycle alternates between a tsetse fly and mammalian host. Tsetse fly host Mammalian host Dividing form (long) Metacyclic form Acquires new VSG Insect bite Saliva Salivary Nervous gland system Changes VSG **Epimastigote** every 1-2 weeks Insect bite Procyclic form Loses VSG Nondividing form (short)



Figure 17.15 VSG genes may be created by duplicative transfer from an internal or telomeric basic copy into an expression site, or by activating a telomeric copy that is already present at a potential expression site. Figure 17.16 Internal basic copies can be activated only by generating a duplication of the gene at an expression-linked site. Initial gene set Probe for basic copy Probe for basic copy in nonexpressing cell in expressing cell Inactive telomeric copy Figure 17.17 Telomeric basic copies can be activated in situ; the size of the restriction fragment Internal basic copy may change (slightly) when the telomere is extended. ELC (active) Basic copy Inactive basic copy Probe for basic copy Probe for basic copy in nonexpressing cell in expressing cell Active telomeric gene Active basic copy Basic copy Activate telomeric copy in situ Copy internal basic gene into expression site Copy telomeric basic gene into expression site



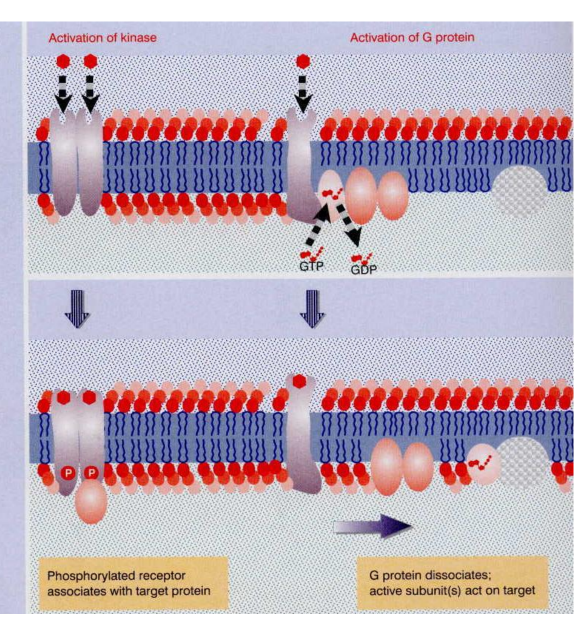


Figure 26.10 Classes of G proteins are distinguished by their effectors and are activated by a variety of transmembrane receptors.

G protein	Effector function	Second messenger	Example of receptor
S	Stimulates adenylyl cyclase	↑ cAMP	β-adrenergic
olf	Stimulates adenylyl cyclase	↑ cAMP	Odorant
	Inhibits adenylate cyclase	↓ cAMP	Somatostatin
	Opens K ⁺ channels	Membrane potential	Somatostatin
0	Closes Ca ²⁺ channels	↓ Membrane potential	m2 acetylcholine
t (transducin)	Stimulates cGMP phosphodiesterase	↓ cGMP	Rhodopsin
q	Activates phospholipase Cb	↑ InsP3, DAG	m1 acetylcholine



Figure 26.3 A signal may be transduced by activating the kinase activity of the cytoplasmic domain of a transmembrane receptor or by dissociating a G protein into subunits that act on target proteins on the membrane.





Locus control region - LCR

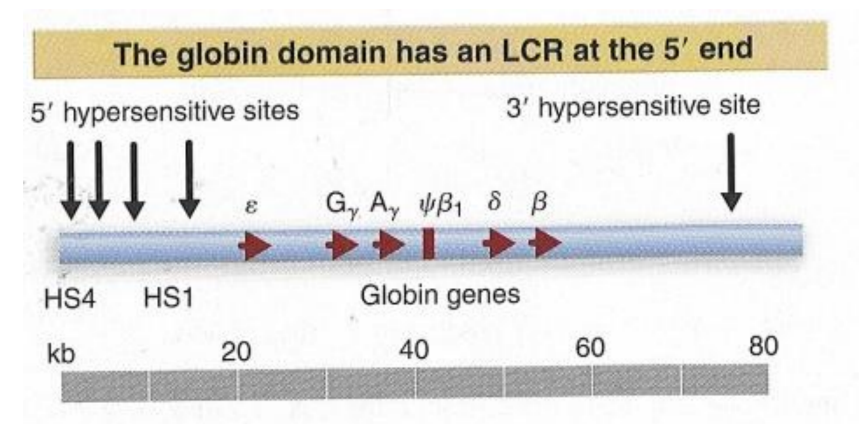


Figure 29.41 A globin domain is marked by hypersensitive sites at either end. The group of sites at the 5' side constitutes the LCR and is essential for the function of all genes in the cluster.



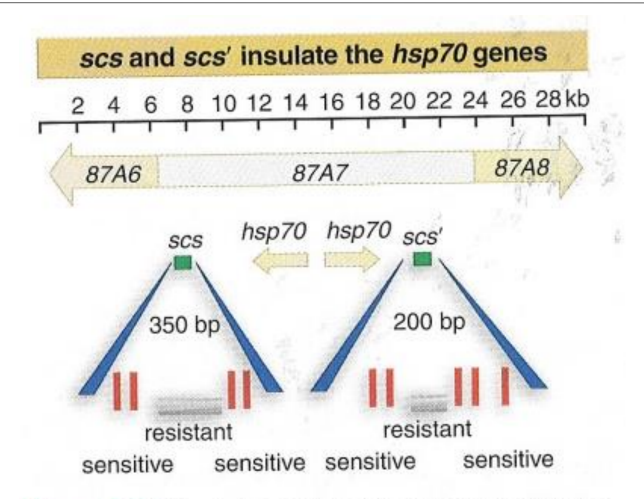


Figure 29.39 Specialized chromatin structures that include hypersensitive sites mark the ends of a domain in the *D. melanogaster* genome and insulate genes between them from the effects of surrounding sequences.



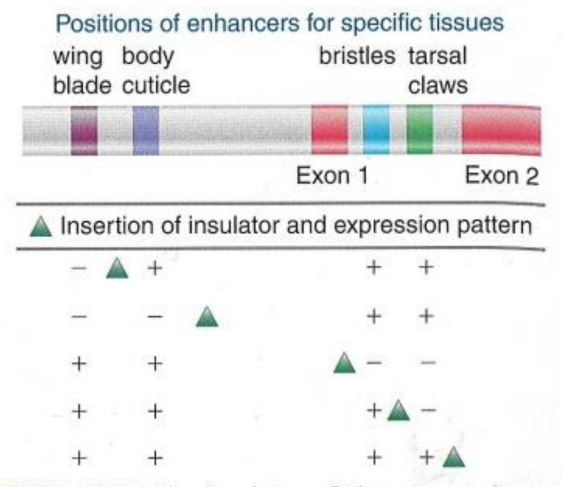
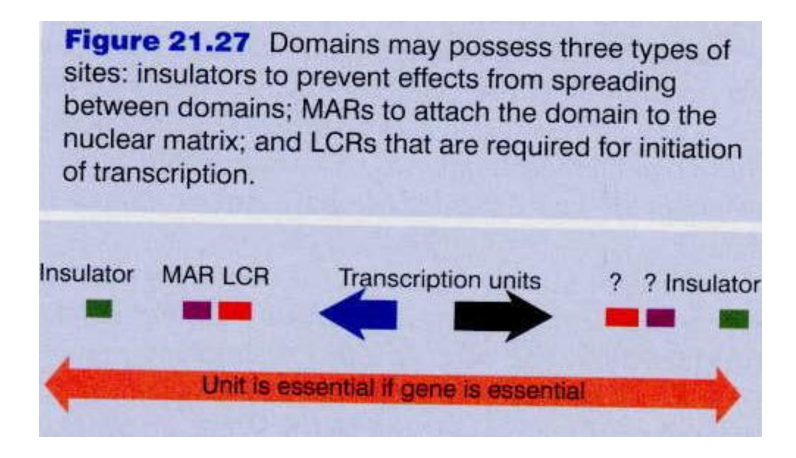


FIGURE 10.54 The insulator of the gypsy transposon blocks the action of an enhancer when it is placed between the enhancer and the promoter.



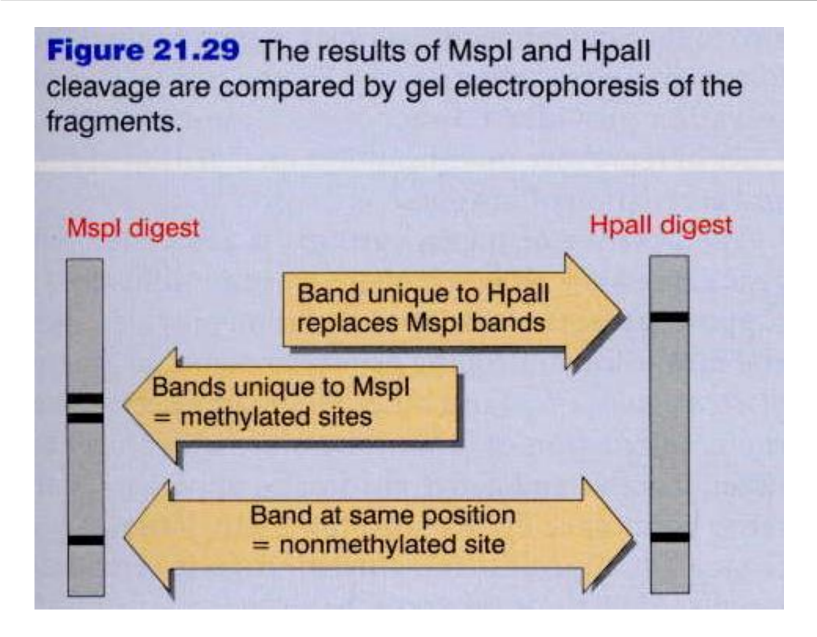


MAR: Matrix attachment site

LCR: Locus control region

Insulator: prevents influence from surrounding regions







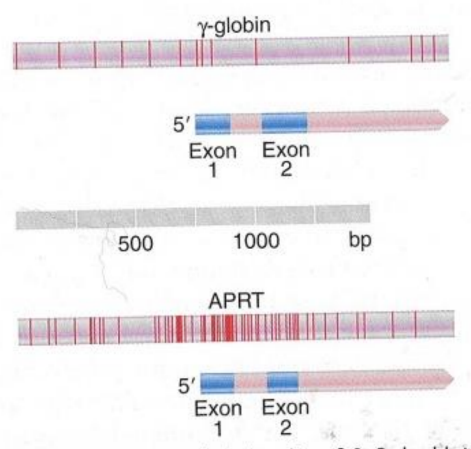


FIGURE 20.18 The typical density of CpG doublets in mammalian DNA is $\sim 1/100$ bp, as seen for a γ -globin gene. In a CpG-rich island, the density is increased to >10 doublets/100 bp. The island in the APRT gene starts ~ 100 bp upstream of the promoter and extends ~ 400 bp into the gene. Each vertical line represents a CpG doublet.



15.12.15