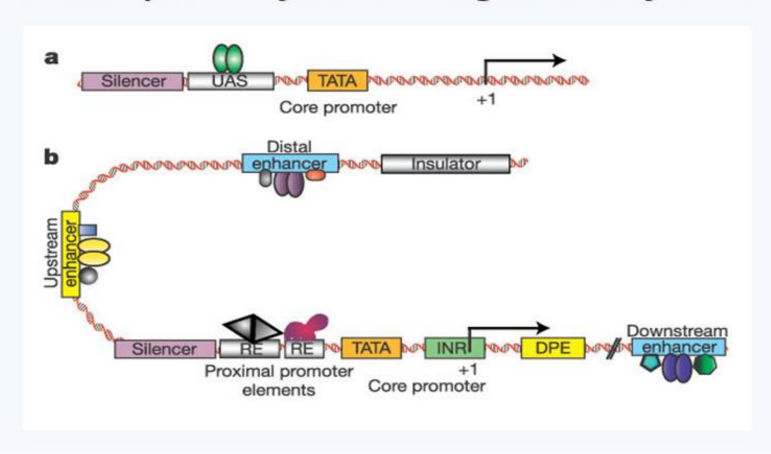
# Transcription activation in multicellular organisms.

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### Regulatory elements controlling RNA Pol. II transcription in yeast and higher eukaryotes.



- Promoters
- Proximal regulatory elements
- Enhancers

# 25.7 Response Elements Are Recognized by Activators

• Response elements may be located in promoters or enhancers. Proximal regulatory elements were targets of the earliest studies.

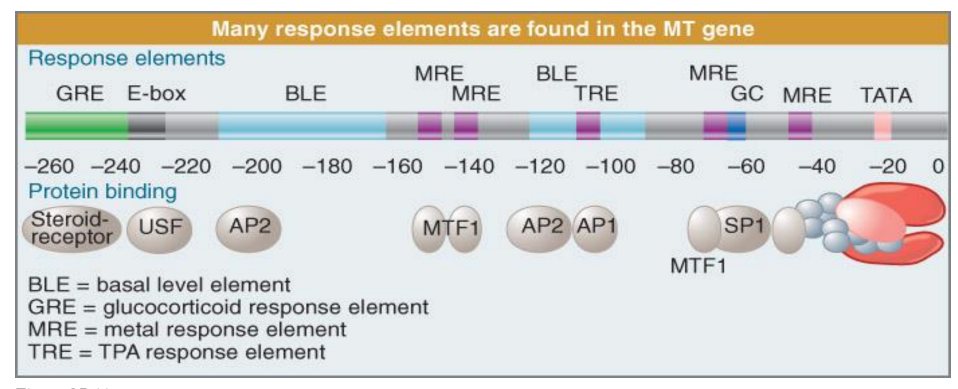
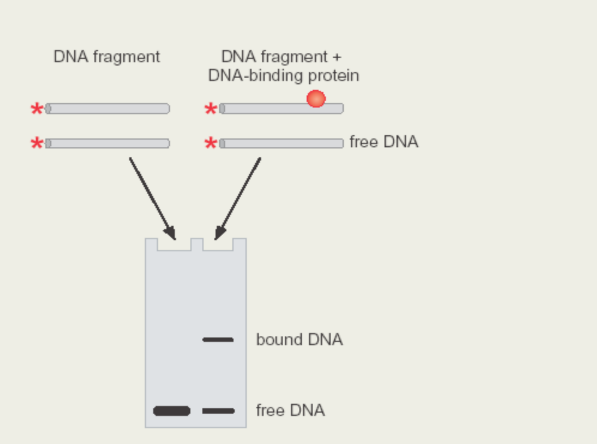


Figure 25.11

#### Many gene regulators from mammalian cells were identified and purified using the Bandshift assay or Gel mobility shift assay.

(also Gel Retardation assay, Electrophoretic Mobility Shift Assay (EMSA)).



shift assay. The principle of the mobility shift assay is shown schematically. A protein is mixed with radiolabeled probe DNA containing a binding site for that protein. The mixture is resolved by acrylamide gel electrophoresis and visualized using autoradiography. DNA not mixed with protein runs as a single band corresponding to the size of the DNA fragment (left lane). In the mixture with the protein, a proportion of the DNA molecules (but not all of them at the concentrations used) binds the DNA molecule. Thus, in the right-hand lane, there is a band corresponding to free DNA, and another corresponding to the DNA fragment in complex with the protein.

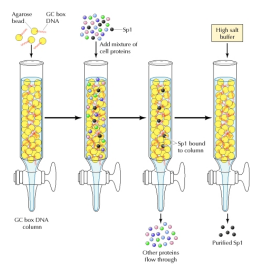
BOX 16-1 FIGURE 2 Gel mobility

Gel is non-denaturing and buffer is mild to avoid disrupting complex. Gel matrix 'traps' protein DNA complex keeping components together to prevent disassociation.

#### Advantages of the bandshift assay.

- DNase1 footprinting required pure proteins but bandshift assays could detect different scarce DNA-binding proteins even in crude nuclear extracts.
- If the probe DNA fragment comes from an enhancer then a point mutation that inactivates the enhancer should also prevent DNA binding by the key regulatory protein in the extract. (Some proteins bind DNA ends or other sites in the probe DNA)
- The specific bandshift provides a convenient assay to purify the regulatory protein.

### DNA-affinity chromatography using a specific oligonucleotide target sequence can purify the binding protein in one step.



Oligonucleotide attached to affinity column contains repeats of a DNA binding site for Sp1 prote

(5'-GGGCGGG-3')N

Figure 6.25 Purification of Sp1 by DNA-affinity chromatography

A double-stranded oligonucleotide containing repeated GC box sequences is bound to agarose beads, which are poured into a column. A mixture of cell proteins containing Sp1 is then applied to the column; because Sp1 specifically binds to the GC box oligonucleotide, it is retained on the column while other proteins flow through. Washing the column with high salt buffer then dissociates Sp1 from the GC box DNA, yielding purified Sp1.

From: Regulation of Transcription in Eukaryotes



The Cell: A Molecular Approach. 2nd edition Cooper GM. Sunderland (MA): Sinauer Associates; 2000

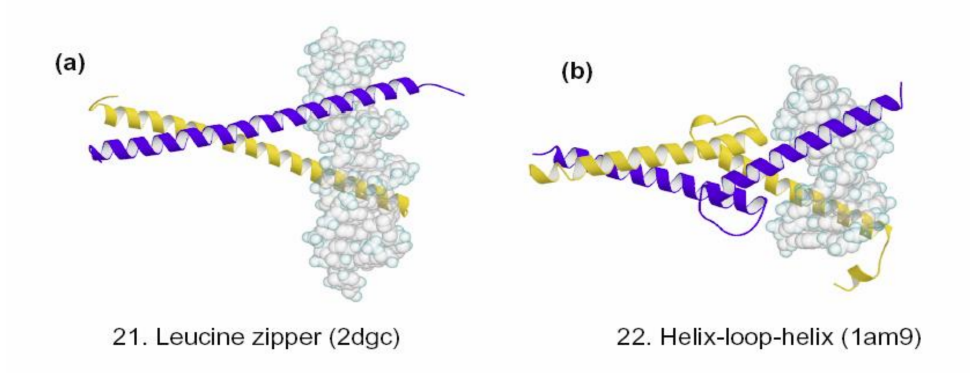
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Examples of important transcription regulators identified after purification from nuclear extracts.

- Fos/jun, AP1. oncoproteins studied by Tom Curran and others. Steve MacKnight showed these proteins contain a leucine zipper.
- Nf-kappaB (Baltimore lab, MIT) Nuclear factor binding the immunoglobulin kappa B enhancer. Important protein in innate immunity and response to viral infection.
- **SRF** (Maniatis lab and Richard Treisman in London) Serum response factor activates genes involved in growth of cells in tissue culture.
- **SP1** (Tjian lab)

### The simplest sequence-specific DNA recognition mechanism of all! CAAT/Enhancer binding protein (C/EBP).



The mechanism of transcription activation by the genespecific regulators.

• The primary idea is the same as for GAL4 in yeast.

• Recruitment of transcription inititation factors and RNA polymerase to the promoter by DNA loop formation between the enhancer and the promoter.

#### Mediator.

- *In vitro* transcription extracts were developed in which Pol II and the General Transcription Factors responded with increased transcription when GAL4 or another activator was added.
- This allowed further purifications to isolate additional components needed to mediate the activation. Depending on what activator was used this purified coactivators or mediator complex.
- Some argue that the complex of Pol II, GTFs and Mediator can exist as a 'holoenzyme' that recruits to promoters as in *E. coli*.

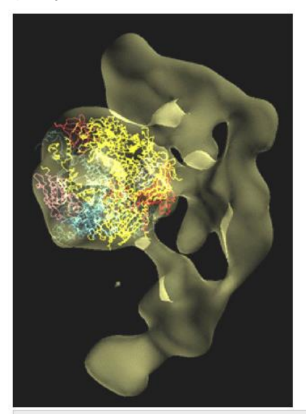
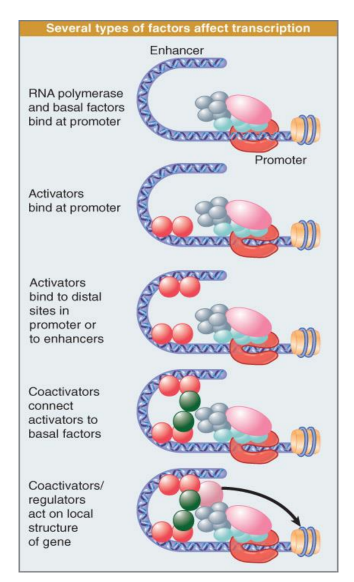


Fig. 18.

Cryo-EM structure of an RNA polymerase II-Mediator complex. The pol II structure was docked in the central density, and is shown in a similar direction of view and color scheme as Fig. 5.

## 25.2 Gene specific transcription factors (Activators/Coactivators) are distinct from the General Transcription factors .



- The basal apparatus determines the startpoint for transcription.
- Activators determine the frequency of transcription.
- Activators work by making protein-protein contacts with the basal factors.
- Activators may work via coactivators.
- Some components of the transcriptional apparatus work by changing chromatin structure. Coactivator p300/CBP (CREB-binding protein) is a histone acetylase targeting H4.

Figure 25.2

• Several factors in the basal apparatus are targets with which activators or coactivators interact. Many activators contact different TAFs. Acidic activators

contact TFIIB **Activators contact the basal apparatus** Activator contacts TAF in TF<sub>II</sub>D Activator contacts TF<sub>II</sub>B

Figure 25.8

• An activator that does not have an activating domain may work by binding a coactivator that has an activating domain. Strong viral activators like Herpesvirus VP16 and Adenovirus E1A are recruited by binding other proteins.

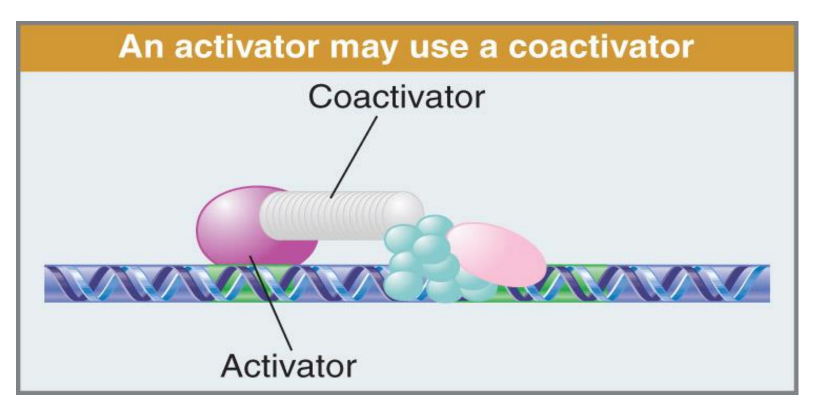
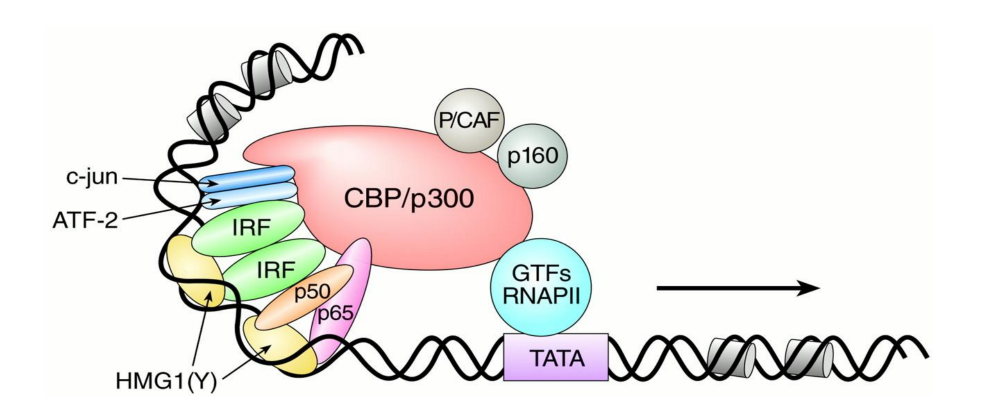


Figure 25.7

#### The IFN- $\beta$ enhanceosome complex.



Vo N, Goodman R H J. Biol. Chem. 2001;276:13505-13508



#### Transcription activation in chromatin.

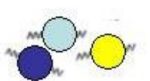
- RNA polymerase was not able to initiate transcription at promoters and some sequence-specific DNA binding proteins could not bind their sites if histones were added first (pioneer transcription factors).
- Getting transcription initiation and later regulated transcription on chromatin templates required extra factors that operate on nucleosomes.
- Constructs with enhancers reintroduced in mammalian chromosomes were often silenced. This turned out later to be due to lack of regulatory elements like the globin gene Locus Control Region (LCR).

### Chromatin immunoprecipitation for genome-wide location analysis of chromosomal proteins in living cells

- 1. Formaldehyde cross link DNA to transcription factors.
- 2. Lyse cells and fragment chromatin by sonication
- 3. Immunoprecipitation



Fragments bound to anti-X antibodies Purify on beads



Unbound fragments are washed away

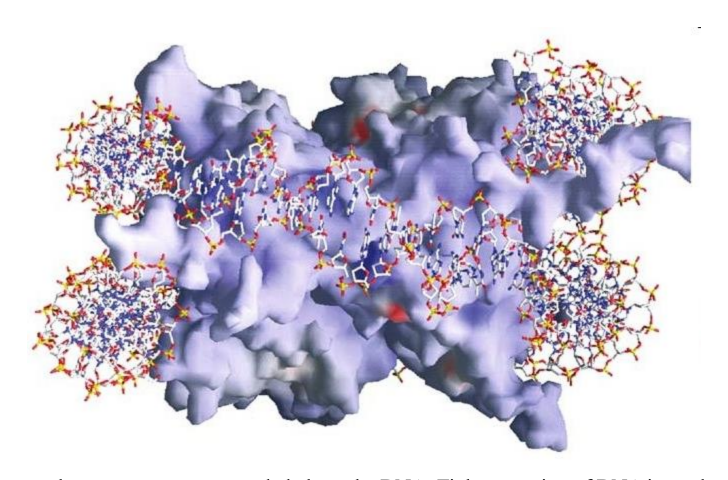
Factor X bound to target gene

- 4. Reverse cross-links, extract DNA
- 5. Analyse which DNA fragments are enriched in the immunoprecipitated fraction.
- **6.** Best done now by Next Generation sequencing (ChIP-Seq)
- 7. Compare reads of immunoprecipitated sequences back to the genome sequence to locate when the protein was bound.

Mapping Polycomb-repressed domains in the bithorax complex using in vivo formaldehyde cross-linked **chromatin**. Orlando V, **Paro** R. Cell. 1993 Dec 17;75(6):1187-98.

There is a contrast between a sequence-specific DNA-binding protein and the DNA packaging nucleosome.

Histone amino acid side chains do not enter the major groove and 'read' the DNA sequence.



• ... but nucleosomes are very crowded along the DNA. Tight wrapping of DNA in nucleosomes means they have positional preferences to bind over more bendable A-T rich sequences first. Very G-C sequences are covered last or not at all.

Most yeast promoters are relatively "nucleosome free" ...many mammalian promoters also....

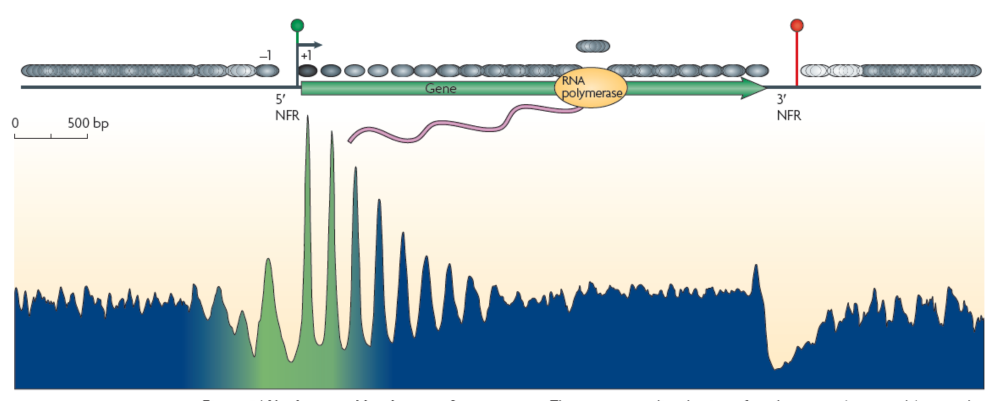


Figure 2 | **Nucleosomal landscape of yeast genes**. The consensus distribution of nucleosomes (grey ovals) around all yeast genes is shown, aligned by the beginning and end of every gene. The resulting two plots were fused in the genic region. The peaks and valleys represent similar positioning relative to the transcription start site (TSS). The arrow under the green circle near the 5' nucleosome-free region (NFR) represents the TSS. The green –blue shading in the plot represents the transitions observed in nucleosome composition and phasing (green represents high H2A.Z levels, acetylation, H3K4 methylation and phasing, whereas blue represents low levels of these modifications). The red circle indicates transcriptional termination within the 3' NFR. Figure is reproduced, with permission, from REF. 20 © (2008) Cold Spring Harbor Laboratory Press.

Relatively nucleosome-free regions, DNAse hypersensitive sites and specific histone modifications help identify enhancers in chromatin.

- Chromatin immunoprecipitation methods are easier to use with abundant nucleosomes than with scarce sequence-specific DNA-binding proteins.
- Modified histones associated with active (H3K4 Me1,Me3) and inactive (H3K27 Me3) genes have been mapped extensively.

  This is a proxy for mapping the regulatory proteins themselves and allows likely enhancers and promoters to be identified.
- Promoters and enhancers also tend to be nucleosome-free.

## Genome-wide locations of the coactivator P300/cEBP identify enhancers.

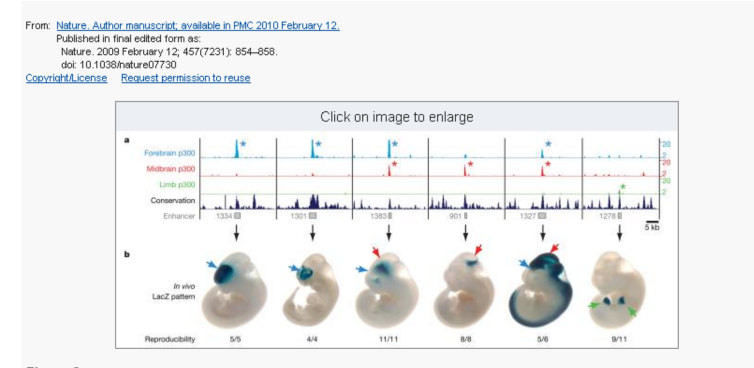


Figure 3

Examples of successful prediction of *in vivo* enhancers by p300 binding in embryonic tissues

**a,** Coverage by extended p300 reads in forebrain (blue), midbrain (red) and limb (green). Asterisks indicate significant (FDR < 0.01) p300 enrichment in chromatin isolated from the respective tissue. Multispecies vertebrate conservation plots (black) were obtained from the UCSC genome browser<sup>50</sup>. Grey boxes correspond to candidate enhancer regions. Numbers at the right indicate overlapping extended reads. b, Representative LacZ-stained embryos with *in vivo* enhancer activity at E11.5. Reproducible staining in forebrain, midbrain and limb is indicated by arrows. Numbers show the reproducibility of LacZ reporter staining. Additional embryos obtained with each construct and genomic coordinates are available using the enhancer ID in the bottom portion of **a** at the Vista Enhancer Browser<sup>32</sup>.

#### Disease mutations in human enhancers.

• Until now human disease mutations in noncoding sequences were very difficult to identify.

• Once enhancers are identified across the human genome they can be included with exons and promoters in focused searches for disease-causing mutations.

#### Pre-mRNA processing in eukaryotes.

- Ribozymes and the RNA world-

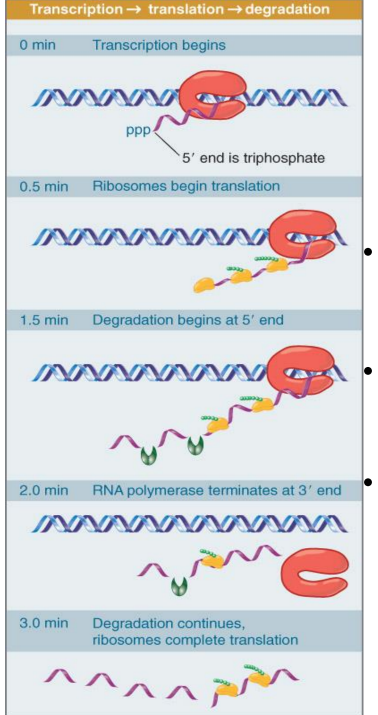
#### Lecture outline

• Eukaryotic mRNA stability, 5' cap and polyA addition and splicing.

• Mechanism of pre-mRNA splicing and alternative splicing. RNA editing.

• Self-splicing Group II introns, RNA catalysis, Ribozymes and the The RNA World.

# Capping and polyA tailing of pre-mRNAs.



#### The life cycle of a bacterial mRNA.

- Bacterial mRNA is unstable and has a half-life of only a few minutes.
- Transcription in both bacteria and eukaryotes is not very fast at 40 nucleotides/sec, 2.4 kb/min.
- Most bacterial mRNAs can be transcribed in a few minutes whereas a human transcript of 100 kb would take 40 minutes.

Figure 7.14

#### hnRNA/pre-mRNA processing

often >1000nt

A typical eukaryotic mRNA:

leader <300nt

•

•	5'cap		AA	.AAAAAAAAn
•	nontranslated	coding	long 3' trailer	n =100-200

## Pre-mRNA 5' cap addition and 3' polyadenylation and splicing are completed before export from the nucleus.

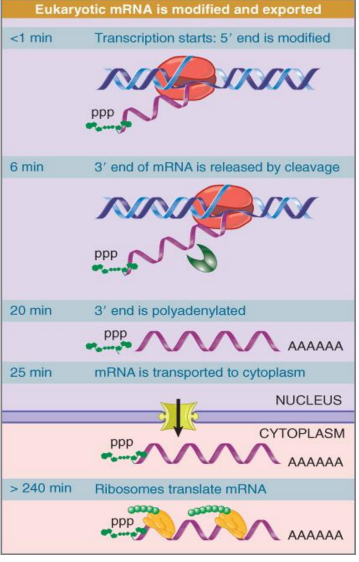


Figure 7.17

# The 5' End of Eukaryotic mRNA Is Capped to protect it against 5' exonucleases.

- A 5' cap is formed by adding a G to the terminal base of the transcript via a 5'-5' link.
- The capping enzyme is **guanylyl transferase.** 
  - 1-3 methyl groups are added to the base of the new terminal guanosine
  - 2 methyl-riboses near cap.
  - Base modification m6A in 5' cap and also inside mRNAs at RRACH consensus, especially around stop codon.

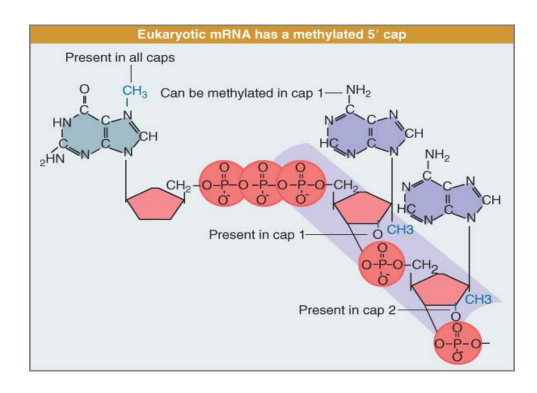
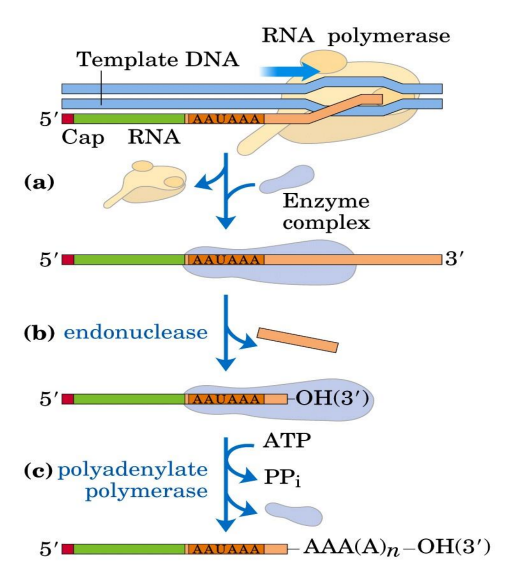


Figure 7.18

#### 3' end cleavage and polyadenylation.



The cleavage and polyadenylation specificity factor (CPSF) recognizes **AAUAAA (AtwoU, Athree)**.

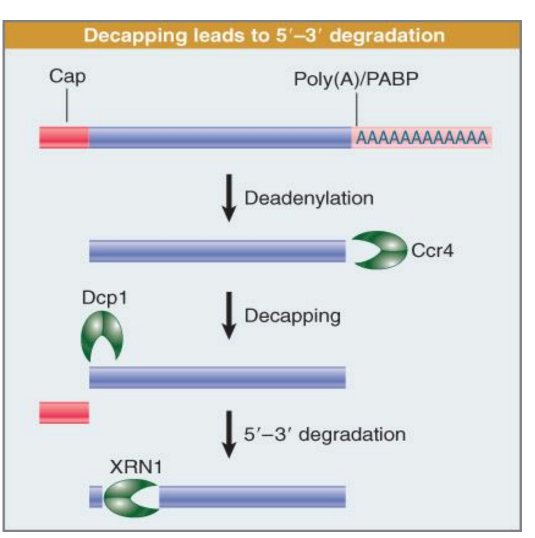
Cleavage factors (CFI and CFII), endonuclease cleaves RNA 30 bases downstream.

Poly(A) polymerase add ~200 A residues processively to the 3' end.

## The polyA tail stabilizes mRNA, facilitates translation.

- Poly A tail is >200 nucleotides long and binds many copies of Poly A-binding protein (PABP).
- PABP stabilizes the mRNA against degradation.
- PABP binds eIF4G (eukaryotic initiation factor 4G) and facilitates translation of message by the ribosome.
- 5' and 3' ends are thought to contact one another by forming a loop of the mRNA.

## Removal of cap and polyA tails are required for regulated degradation of mRNA.



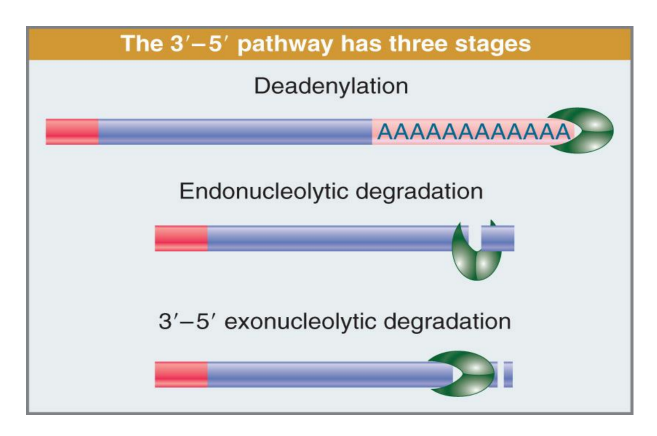
• Degradation of yeast mRNA requires removal of the 5' cap and the 3' poly(A).

 One yeast pathway involves exonucleolytic degradation from 5'→3'.

Figure 7.24

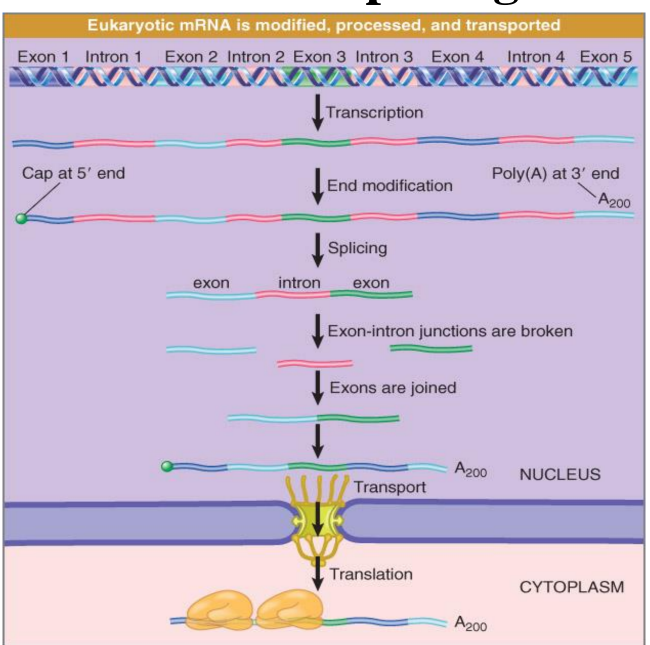
• Another yeast mRNA degradation pathway uses a complex of several exonucleases that work in the  $3' \rightarrow 5'$  direction The **exosome**.

• The deadenylase of animal cells may bind directly to the 5' cap.

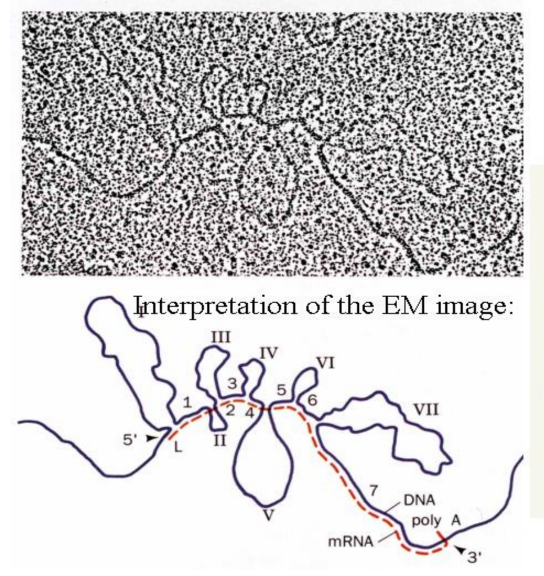


### The mechanism of eukaryotic premRNA splicing.

#### Pre-mRNA splicing.

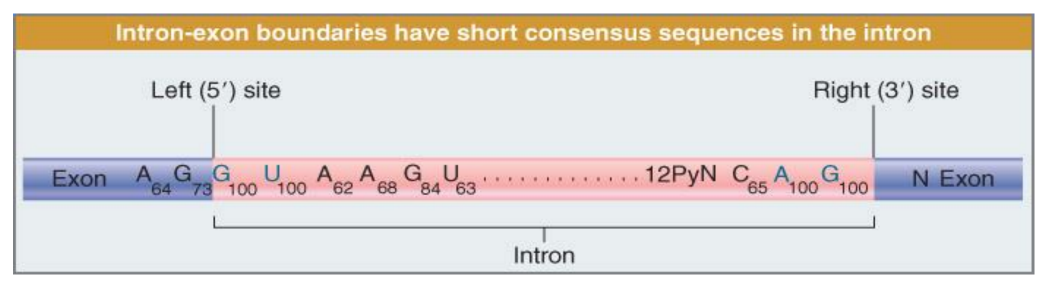


## EM heteroduplex analysis of Adenovirus RNA transcripts annealed to Adenovirus DNA showed formation of D (displacement) loops.



Annealing RNA from virus-infected cells with viral DNA revealed the existence of seven introns-transcribed regions of the DNA removed from the mature mRNA.

## Consensus 5' and 3' splice sites. The GU-AG rule



Protein coding sequence must be 'in frame' across the splice junction.

Remember GU...AG are splice junction sequences. Mnemonic; GULAG maybe?

Intron sequences are generally not conserved even between closely related genes. Figure 26.3

# Splicing must preserve the open reading frame.

• 5' and 3' splice site have to be in the same reading frame.

• Reading frames are 1, after first base of codon, 2 after second base and 3 after third base, ie. between codons

• GU..AG consensus are the ends of the intron to be removed, not in the actual coding sequence.

#### **Splice Junctions Are Read in Pairs**

- All 5' splice sites are functionally equivalent, and all 3' splice sites are functionally equivalent.
- Usually draw exons as boxes to emphasise them with splices joining box corners and introns as lines between the boxes .
- Gene rearrangements within big introns have a good 1/3 chance to produce fusion proteins. (Exon shuffling in evolution).

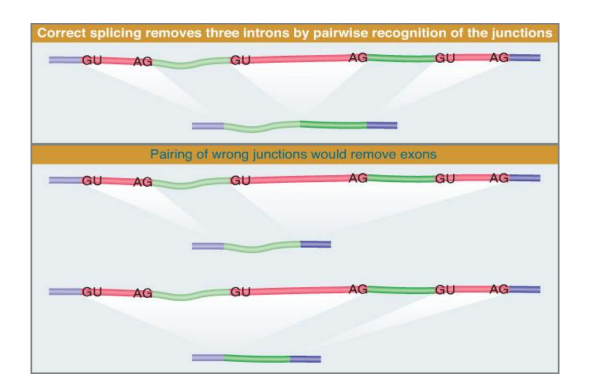
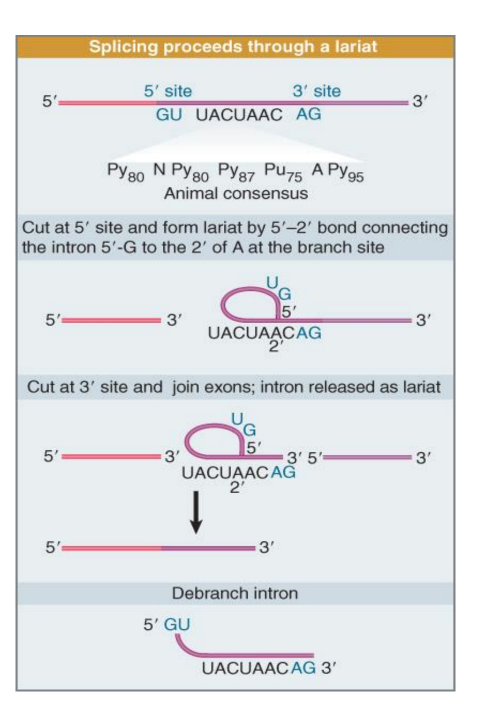
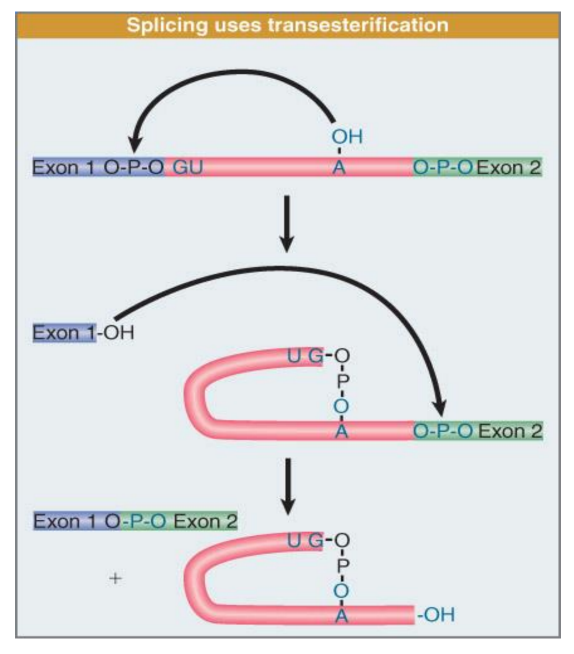


Figure 26.4

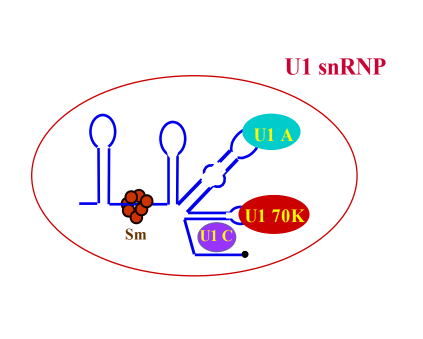


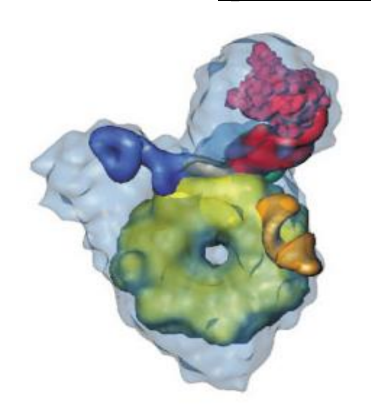


#### Alkaline lysis of ssRNA.

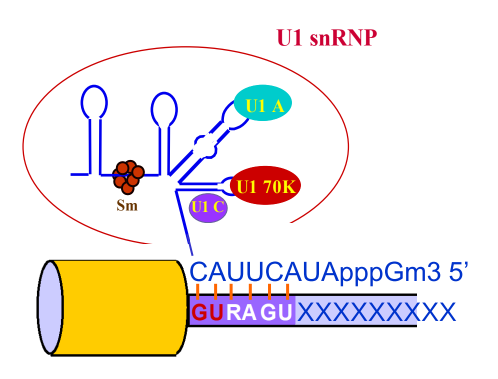
The reactive 2' OH on ribose that mediates alkaline self-cleavage of single stranded RNA Splicing is catalysed by a series of five small nuclear ribonucleoprotein particles (snRNPs, 'SNURPs'). Conserved small structured RNAs with Sm and other proteins bound.

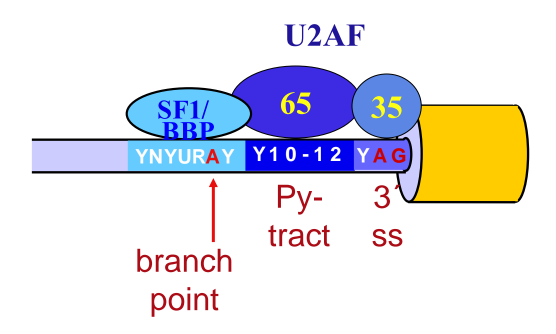
SNURPs assemble sequentially on pre-mRNA to form a spliceosome.

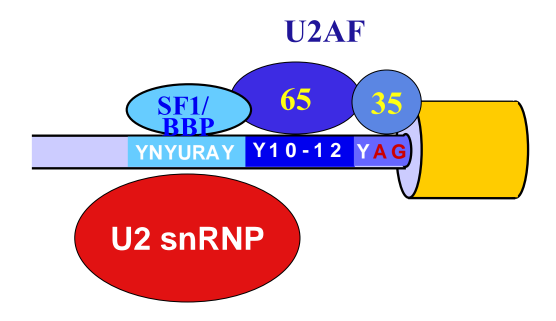


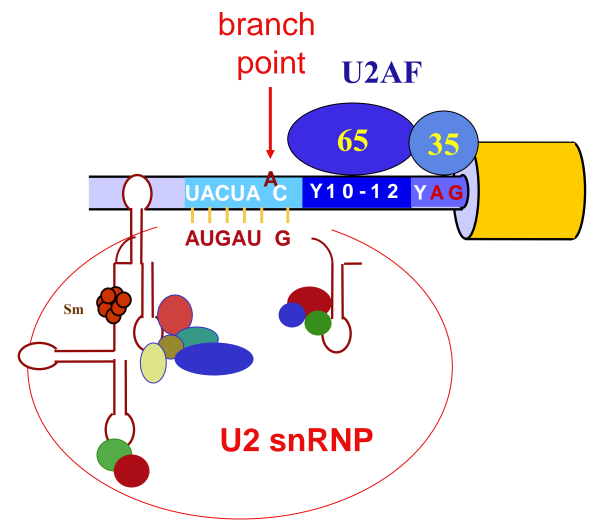


Cryo-EM picture (Stark et al. Nature 409, 539)

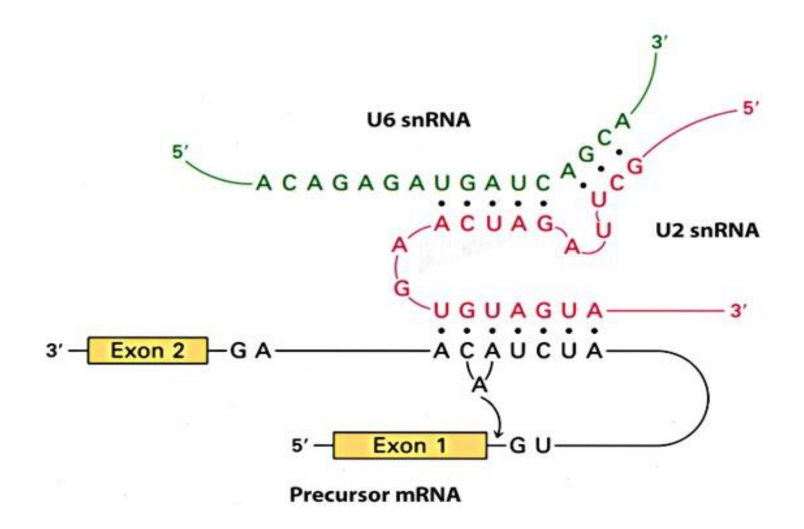




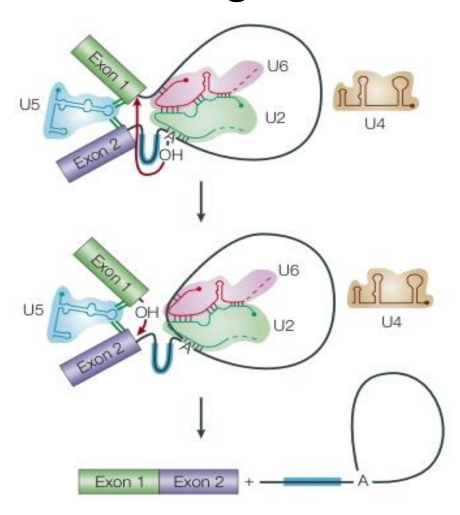




The U4/U5/U6 tri snRNP arrives last and base pairs to U2.



## Conformation changes bring splice junctions together.

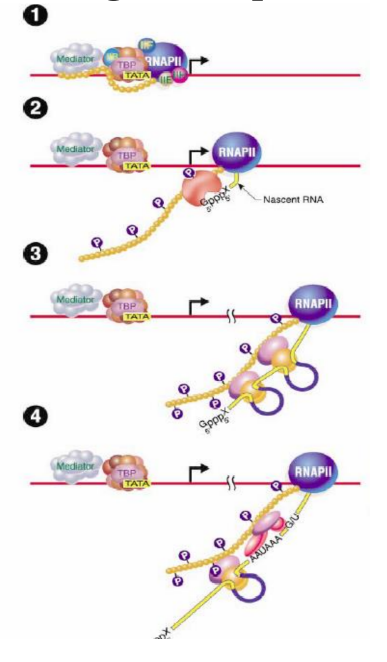


#### Pre-mRNA



spliceosome

RNA processing is coupled to transcription.



## Splicing Is Connected to Export and efficient translation of mRNA

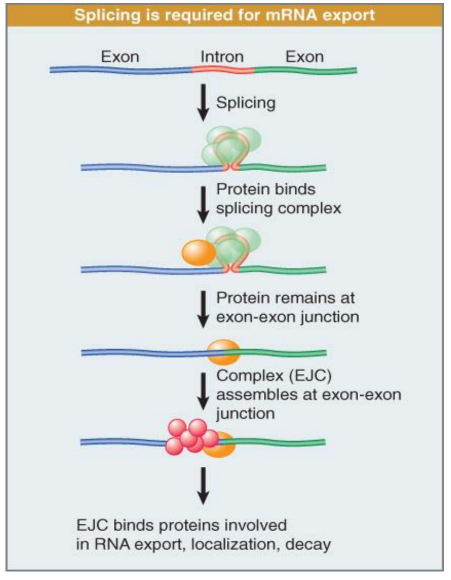
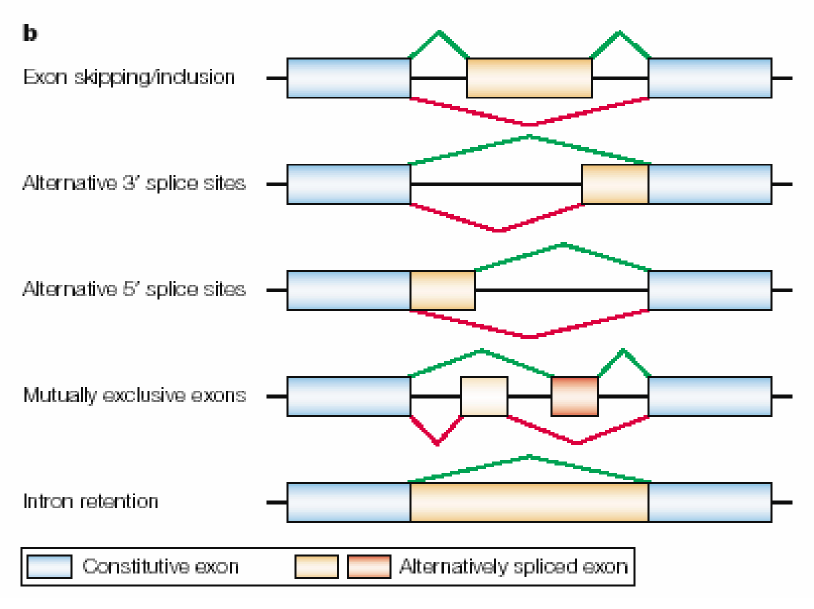


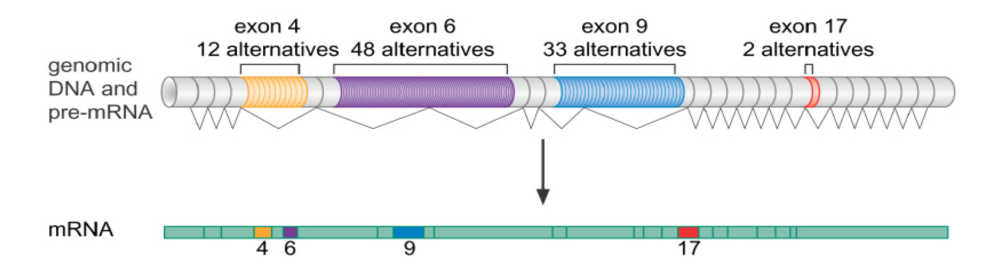
Figure 26.16

Alternative splicing is an important strategy to increase the number of gene products produced from a single gene.

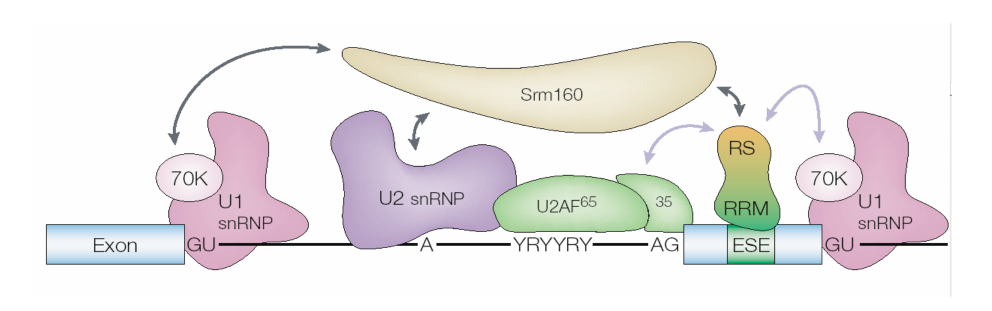
#### Modes of alternative splicing.



### Alternative splicing in *Drosophila* Dscam.



Inclusion of alternative exons may be promoted by exonic splicing enhancers that bind serine/arginine-rich (SR) RNA-binding proteins.



Heterogeneous ribonuclear proteins (hnRNP proteins) coat pre-mRNAs and mRNAs and often inhibit splicing events that SR proteins promote. 'Ying-yang' balance of these effects on many splicing events.

#### Splicing complexes Can be Formed by Intron Definition or Exon Definition

- The direct, **intron definition**, way of forming an E complex at short introns is for:
  - U1 snRNP to bind at the 5' splice site
  - U2AF to bind at a pyrimidine tract between the branch site and the 3' splice site

- Another possibility, **exon definition**, for very long introns is for the complex to form between:
  - U2AF at the pyrimidine tract
  - U1 snRNP at a downstream 5' splice site

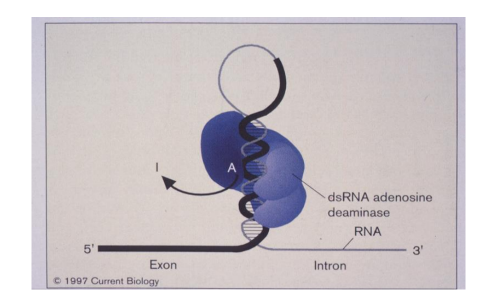
### The importance of alternative splicing

It is estimated that 95% of human genes are alternatively spliced from new sequencing information.

Alternative splicing creates protein diversity 26,000 human protein-encoding genes, yet 90,000 proteins

Aberrant splicing is a frequent effect of human disease gene mutations.

### Base deamination RNA editing also occurs site-specifically in some transcripts and changes codon meaning.



Adenosine deaminases acting on RNA (ADARs) edit 4% of *Drosophila* transcripts (972 sites), and ~250 known sites in human transcripts, including those encoding brain ion-channel subunits, particularly GluR B, the dominant subunit of AMPA class glutamate receptors.

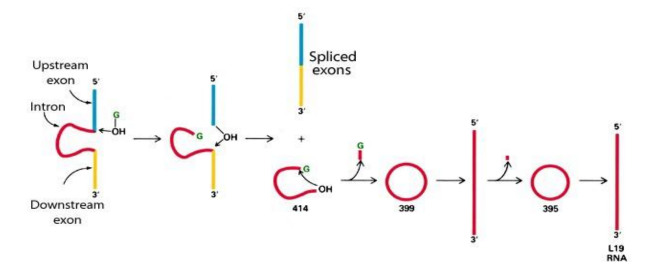
APOBECs are C to U deaminases first identified in RNA ediding but more important as DNA editors. Activation induced deaminase (AID) edits DNA in immunoglobulin genes to initiate Somatic Hypermutation of antibody variable regions. APOBEC3G edits HIV and interferes with virus replication.

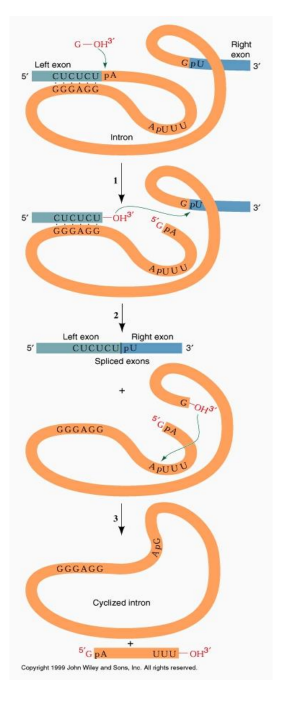
# RNA catalysis, Ribozymes and the RNA World.

#### The RNA World.

Self-splcing intorns are small transposable elements that remove themselves cleanly from the transcripts they are inserted in.

Group I intron splicing in 26S rRNA of *Tetrahymena* is catalysed by RNA in the absence of protein (Tom Cech).



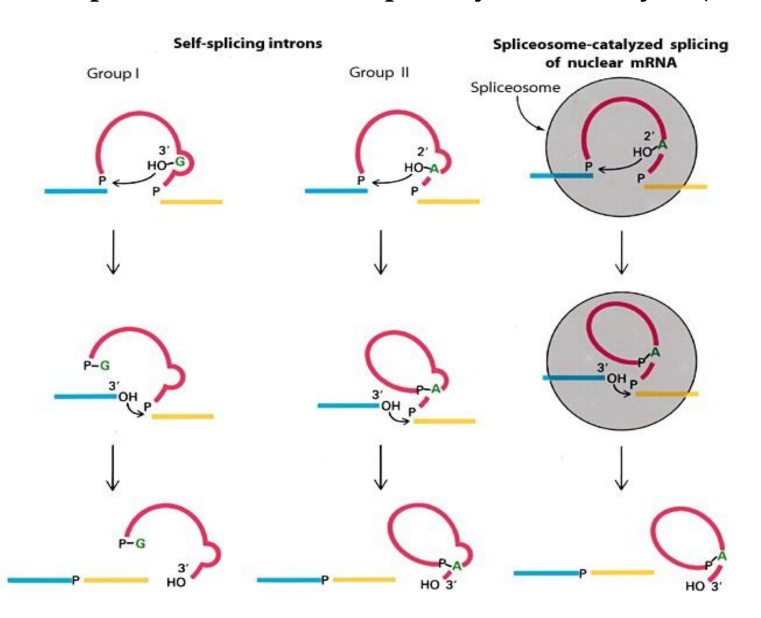


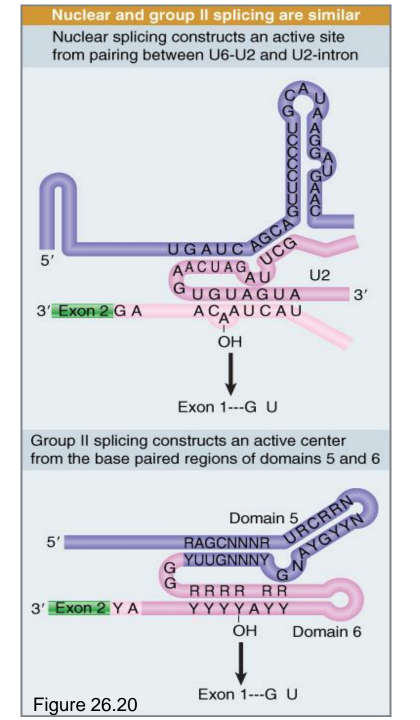
#### The RNA World idea - RIBOZYMES.

- Ribosomes and spliceosomes function through RNA-RNA interactions. Before the self-splicing introns were discovered it was assumed that catalysis would be caused by the proteins in these complexes.
- The discovery that RNA alone is catalytic in the Group I self-splicing intron showed that the RNA could be catalytic in ribosomes and spliceosomes also. RNA enzymes are called **RIBOZYMES** and many artificial ones have been created.
- This contributes to our understanding of possible origins of life as RNA alone could have carried out its own replication using natural ribozyme activities until translation was invented as a way to make more versatile protein catalysts.
- DNA also may have been a later invention for use as a genomic library. It is more stable to hydrolytic cleavage than RNA and more readable by proteins than dsRNA to allow gene control.

The RNA World.

Group II introns also self-splice by RNA catalysis (Cech).





### How introns might have spread. Mobile Group II introns.

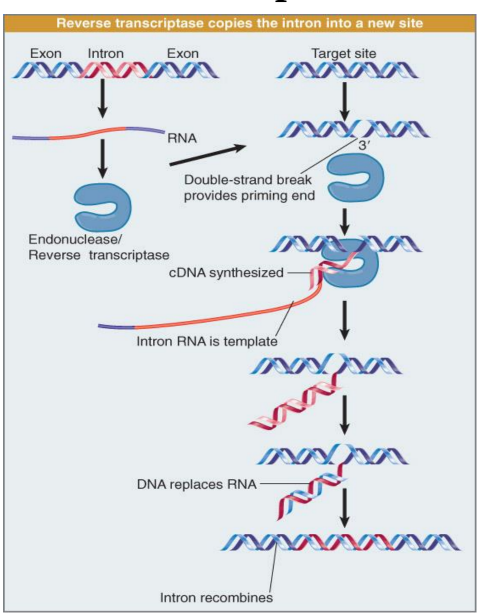
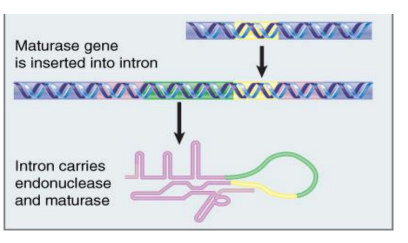


Figure 27.13

# How spliceosomal introns might have evolved from self-splicing Group II introns.



- Autosplicing introns may require maturase protein activities encoded within the intron.
  - They assist folding into the active catalytic structure.
- Imagine that such an intron loses its maturase but can use maturase provided by other similar introns.
- This intron might eventually evolve to consist of just the target splice junctions GU..AG with all the catalytic activities supplied by a trans-acting splicesome.

### Summary.

• Eukaryotic mRNA stability, 5' cap and polyA addition and splicing.

• Mechanism of pre-mRNA splicing and alternative splicing. RNA editing.

• Self-splicing Group II introns, RNA catalysis, Ribozymes and the The RNA World.