

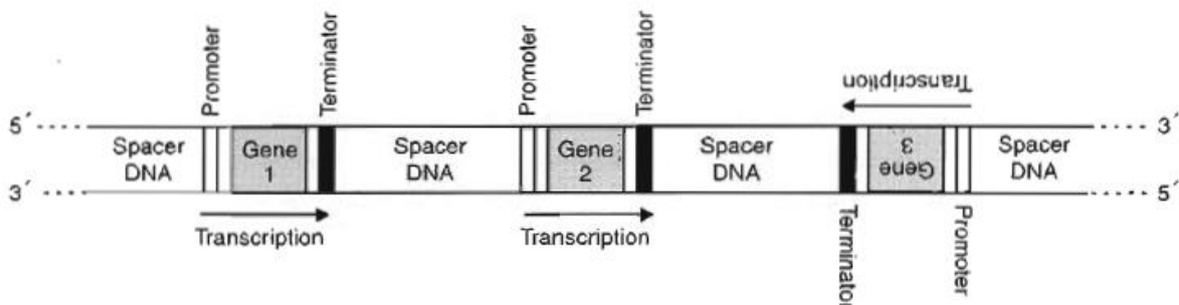
- ✚ Some notes regarding the picture above: (ادرس الصورة مع الملاحظات مليح) 😊
- ✓ Central dogma of life means the principle of life (عقيدة الحياة ومبدأها)
 - ✓ DNA-dependent DNA polymerase means it uses the DNA and its bases sequence to make a DNA molecule
 - ✓ The same for the two other enzymes, RNA-dependent DNA polymerase it uses the RNA to make DNA , DNA-dependent RNA polymerase uses DNA to make RNA
 - ✓ the flow of genetic information happens by two main processes, first, Perpetuation (which means preservation) of genetic information from generation to the next which happens through **replication of the DNA** and the enzyme responsible called **DNA-dependent DNA polymerase**, second, control of the phenotype (المظهر الخارجي) and gene expression through **transcription** and the enzyme her is **DNA-dependent RNA polymerase** and after that the resulting mRNA goes through the process of **translation** to make protein
 - ✓ **Reverse transcription** uses **RNA-dependent DNA polymerase**, this process happens in our body in the activity of the **telomerase** enzyme which depends on short segments of RNA to synthesize the repeated sequences at the end of the DNA molecule, another example is a group of viruses called **retroviruses (has RNA as genetic material)**
 - ✓ **Translation** or protein synthesis happens when the mRNA (result of transcription) leaves the nucleus to be translated on the protein synthesizing machinery which make mature active functional protein (the protein when it comes out from the ribosome it isn't functional protein and it goes through **post translational modifications** to be functional which takes place in **the lemon endoplasmic reticulum and Golgi apparatus** and after that it will be secreted form Golgi by vesicles)

- ✓ The new DNA molecule goes through **post replication modifications**.
- ✓ The mRNA goes through **post transcriptional modifications**.
- ✓ Gene expression means transferring the genetic information from DNA to mRNA then the translation by the ribosome to make protein which means it's a two processes first mRNA transcription and second protein synthesis.

Overview of transcription:

- ✓ The first stage in the expression of genetic information is transcription of the information in the DNA (deoxyribonucleotides sequence) into RNA (ribonucleotides Sequence).
- ✓ For any gene, only one strand of the DNA molecule, called **the template strand**, is transcribed by **RNA polymerase**.
- ✓ Because RNA polymerase moves **in the 3' to 5' direction along the template strand** of DNA, the RNA product is **antiparallel (it'll be 5' to 3')** and **complementary to the template**.
- ✓ The DNA has two strands: **templet non-coding stand** and **non-templet coding strand** (**templet**: go into transcription + its sequence complementary to mRNA/**coding**: doesn't go into transcription + its sequence is the same as the mRNA with the change of thymine and uracil)
- ✓ RNA polymerase is a blind enzyme because it can't recognizes start signals (**promoters**) and stop signals (**terminators**) for any of the thousands of transcription units in the genome of an organism by its own so it needs the help of transcription factors to recognize them.

Transcription of several genes on a chromosome



Types of RNA (RNA molecules play a variety of roles in the cell):

1. Ribosomal RNA (**rRNA**), which is the most abundant (85%) type of RNA in the cell, it is complexed with different type of protein in the form of ribosome (number of protein varies with the type of *ribosome –in prokaryotic or eukaryotic cell-*)
2. Transfer RNA (**tRNA**), which is the second most abundant type of RNA and (it contains thymine + other unusual ribonucleotides), it's the smallest type of RNA (t for tiny) 75-94 ribonucleotides, there are at least 20 type* of tRNA because each one can carry one amino acid to the site of protein synthesis
 *(every type of tRNA can carry only specific amino acid, but there are some amino acid that has more than one tRNA/ so their number is more than 20)
3. Messenger RNA (**mRNA**) the only type of RNA that is translated, which carries *the genetic information from particular gene that determines(SPECIFYING) the amino acid sequence of a particular protein* to the ribosome.

**The mRNA population in a cell is very heterogeneous (مختلف) in size and base sequence, as the cell has essentially a different mRNA molecule for each of the thousands of different proteins made by that cell.

4. Heterogeneous nuclear RNA (**hnRNA or pre-mRNA**) (the immediate(first) product of gene transcription), which is found only in the nucleus of eukaryotic cells and it represents precursors of mRNA, it's immature and inactive so that it needs processing and 75% is degraded in the nucleus and 25% only is processed to mature RNA.
5. Small nuclear RNA (**snRNA**)(ribozymes) (RNA molecules with enzymatic activity), small fragments of RNA 90-300 ribonucleotides, which is also only found in the nucleus of eukaryotes, small in size and complexed with proteins (forming ribonucleoproteins), One of its major functions is to participate in splicing (removal of) intron to make mRNA (it helps the immature hnRNA{that has things called exons and introns} to be mature mRNA by removing the introns..
6. **Micro-RNA**,(formed by enzymes in the nucleus that cut the mRNA), short, non-coding, ~ 22 nucleotide long, *generated by nucleolytic processing* of the products of distinct genes or transcription unites, at least some of which control the expression of other genes during development (mature micro RNA molecules can hybridize together(بتندمج) to form **imperfect** RNA-RNA duplex within the 3' untranslated regions of specific target mRNA causing unexplained gene expression regulation in at least half (50%) of the human genes),its function to regulate the rate of transcription of particular gene.
7. Small cytoplasmic RNA (**scRNA**), has catalytic activity in tRNA processing and acts as signal recognition particle.
8. Small nucleolar (**snoRNA**) acts in rRNA processing/maturation/methylation.
9. Small interfering RNA (**siRNA**) are derived by specific nucleolytic cleavage of larger **double stranded RNAs** to form small 21-25 long products, its function to regulate the rate of transcription of particular gene same as micro-RNA.
 - They form **perfect** RNA-RNA hybrids with their targets(mRNA) anywhere within the length RNA where the complementary sequence exists resulting in reduction of specific protein production because siRNA-mRNA complexes are degraded by nucleolytic machinery (interferes with the expression of specific gene by hybridizing to its corresponding RNA sequence in the target mRNA, then activates degradation of mRNA which can't be translated into proteins).

✚ Transcription: important concepts and terminology

- ✓ RNA polymerase locates genes in DNA by searching for promoter regions.
- ✓ The promoter is the binding site for transcription factors and RNA polymerase.
- ✓ Binding establishes where transcription begins, which strand of DNA is used as the template, and in which direction transcription proceeds.
- ✓ RNA polymerase moves along the template strand in the 3' to 5' direction as it synthesizes the RNA product in the 5' to 3' direction using NTPs (ATP, GTP, CTP, UTP) as substrates.
- ✓ RNA polymerase does not proofread its work.
- ✓ The RNA product is **complementary** and **antiparallel to the template strand**.
- ✓ The coding (**non-template**) strand is not used during transcription. It is identical in sequence to the RNA molecule, except that RNA contains uracil instead of the thymine found in DNA.

- ✓ By convention, the base sequence of a gene is given from the coding strand (5' → 3').
- ✓ Transcription ends when RNA polymerase reaches a termination signal.

RNA Polymerases (prokaryotic cells):

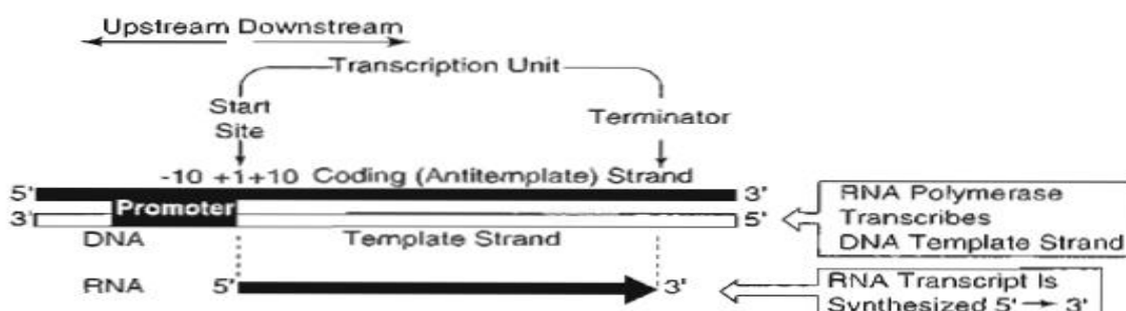
- ✓ There is a single prokaryotic RNA polymerase that synthesizes all types of RNA in the cell. (remember it's a blind enzyme doesn't know where to start or to stop so it needs transcriptional factor to help)
- ✓ The core polymerase has the subunit structure ($\alpha 2\beta\beta'$) (it has four polypeptide chains).
- ✓ RNA polymerase ($\alpha 2\beta\beta'$) + sigma (σ) factor = holoenzyme
- ✓ A protein factor called sigma (σ) is required for the initiation of transcription at the promoter.
- ✓ σ factor is released immediately after transcription initiation.
- ✓ **Functions of the subunits:**
 - α : assembly of the tetrameric core (الي يتربط الاربع وحدات مع بعض)
 - β : ribonucleoside triphosphate binding site, link ribonucleotides together with (phosphodiester bond)
 - β' : DNA template binding region
- ✓ **σ (sigma factor):** initiation of transcription
- ✓ The transcription can be ended by a particular sequence on the mRNA or
- ✓ Termination of transcription sometimes requires a protein called rho (ρ) factor.
- ✓ This enzyme is inhibited by rifampin and actinomycin D.

Promoter "Strength" (activity):

- ✓ What determines the strength of promoter is the number of initiation of transcription per unit time.
- ✓ Affects amount of RNA made, so, it affects level (rate) of expression for that gene.
- ✓ Not all promoters have same "strength"
- ✓ Promoters differ in DNA sequences and "strength"
- ✓ RNA polymerase binds differently to different sequences / "Strong promoters" initiate transcription more often than "weak promoters"
 - rRNA has strong promoter: ~1 initiation per second
 - lacZ (اشي بيهضم اللاكتوز بالبكتيريا) has a weak promoter: ~1 initiation per minute

Eukaryotic RNA polymerases: there are three types which can be distinguished by the particular types of RNA they produce:

- 1- **RNA polymerase I** (the most specified one) is located in the nucleolus and synthesizes 28S, 18S, and 5.8S rRNAs. (s for sedimentation rate)
- 2- **RNA polymerase II** is located in the nucleoplasm and synthesizes hnRNA/mRNA and some snRNA.



3- **RNA polymerase III** (the least specified one) is located in the nucleoplasm and synthesizes tRNA, some snRNA, and 5S rRNA.

some notes:

- Transcription factors (such as TFIID for RNA polymerase II) help to initiate transcription.
- The requirements for termination of transcription in eukaryotes are not well understood.
- In addition, RNA polymerase II is inhibited by (α -amanitin) a toxin from certain mushrooms. It inactivates RNA pol II and can kill a person, while, RNA pol I and III are less affected by toxin

Comparison of eukaryotic and prokaryotic RNA polymerases

Prokaryotic	Eukaryotic
Single RNA polymerase ($\alpha_2\beta\beta'$)	RNAP 1: rRNA (nucleolus), except 5S rRNA RNAP 2: hnRNA/mRNA and some snRNA RNAP 3: tRNA, 5S rRNA
Requires sigma (σ) to initiate at a promoter	No sigma, but transcription factors (TFIID) bind before RNA polymerase
Sometimes requires rho (ρ) to terminate	No rho required
Inhibited by rifampin \rightarrow Actinomycin D	RNAP 2 inhibited by α -amanitin (mushrooms) Actinomycin D

The following events occur during the expression of a **prokaryotic gene**:

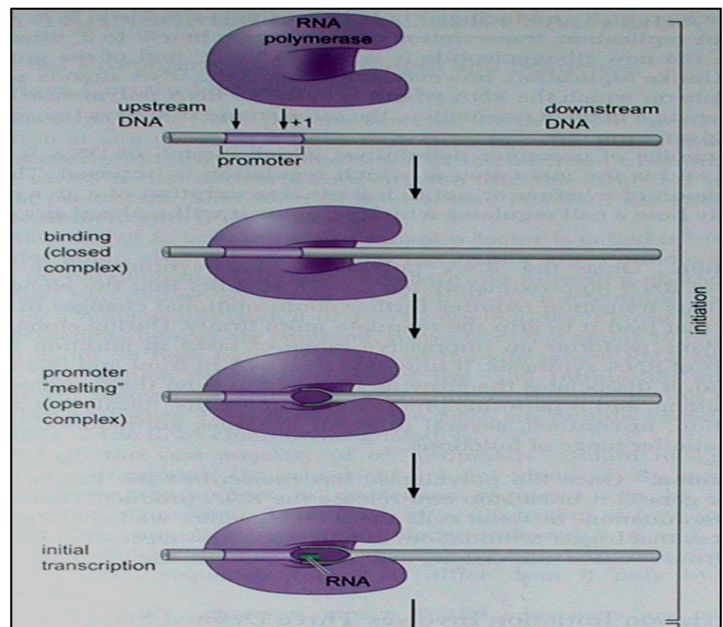
- ✓ With the help of sigma factor, RNA polymerase recognizes and binds to the promoter region.
- ✓ The bacterial promoter contains two "consensus" sequences, called **the Pribnow box** (it lays 10 nucleotides far from the initiation site **-10**) [TATA (TATTAT) box] and the **-35 sequence** (TGTTGACA).
 - 1- The promoter identifies the start site for transcription and orients the enzyme on the template strand.
 - 2- Transcription begins at the + 1 base pair, Sigma factor is released as soon as transcription is initiated.
 - 3- The core polymerase continues moving along the template strand in the 3' to 5' direction, synthesizing the mRNA in the 5' to 3' direction.

Initiation

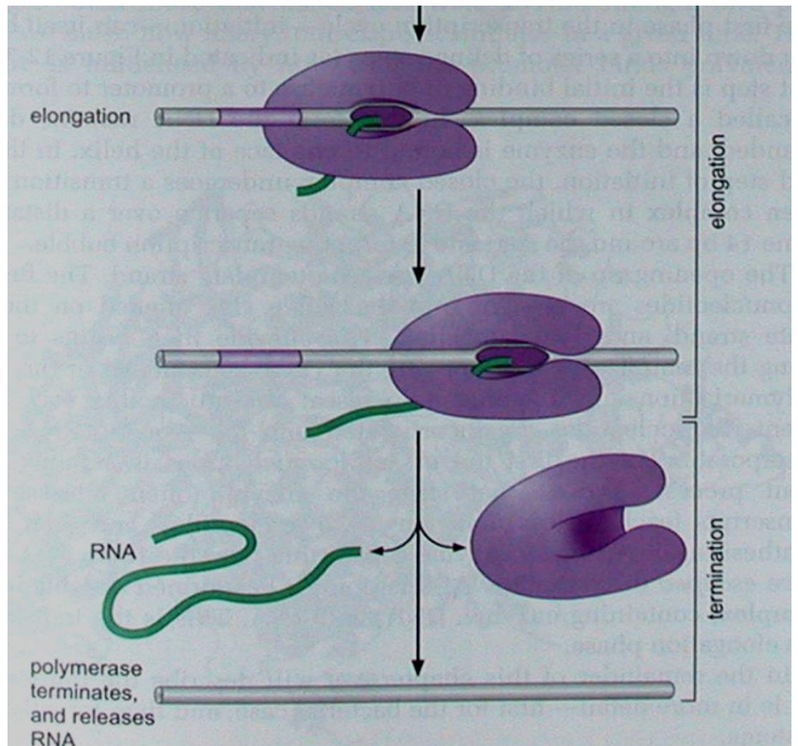
A- Binding (closed complex)

B- Promoter "melting" (open complex)
Binding of the RNA polymerase seems to stimulates the unbinding of the two strand so it can begin its work

C- Initial transcription



Elongation



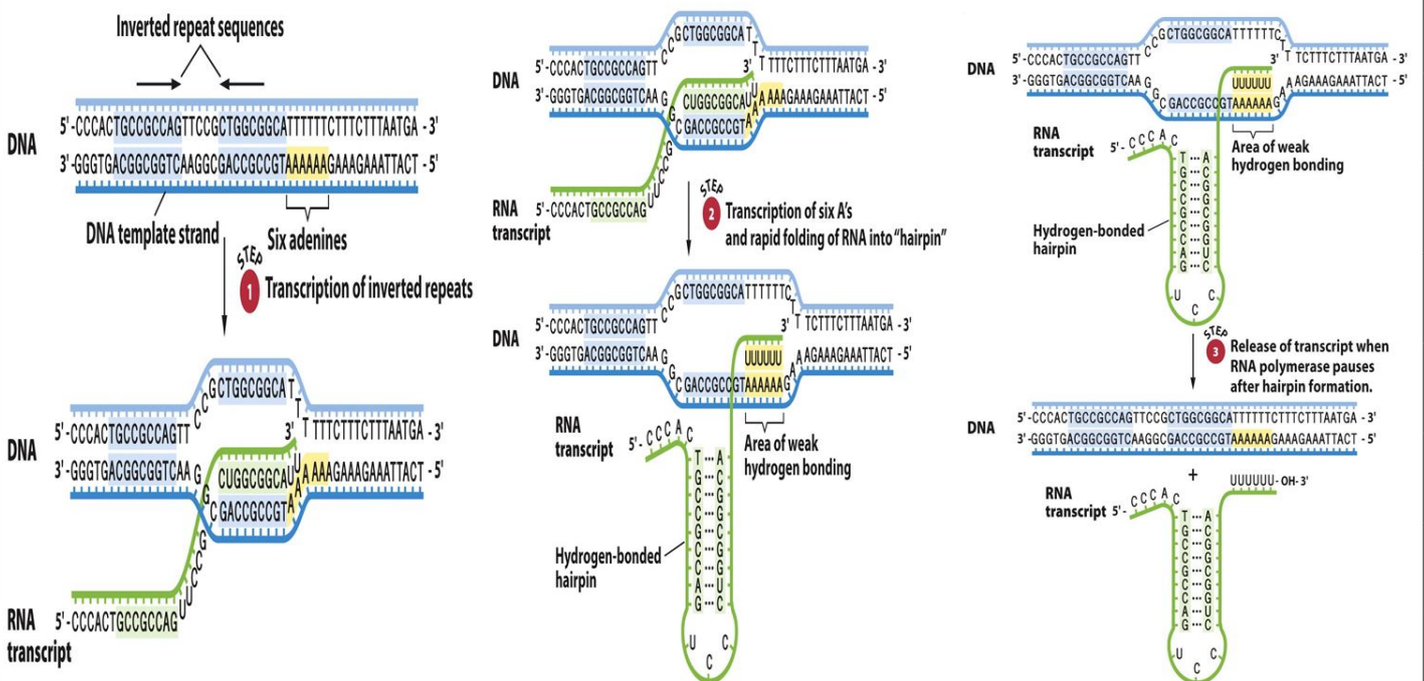
Termination

4- RNA polymerase eventually reaches a transcription termination signal, at which point it will stop transcription and release the completed mRNA molecule.

✓ There are two kinds of transcription terminators commonly found in prokaryotic genes:

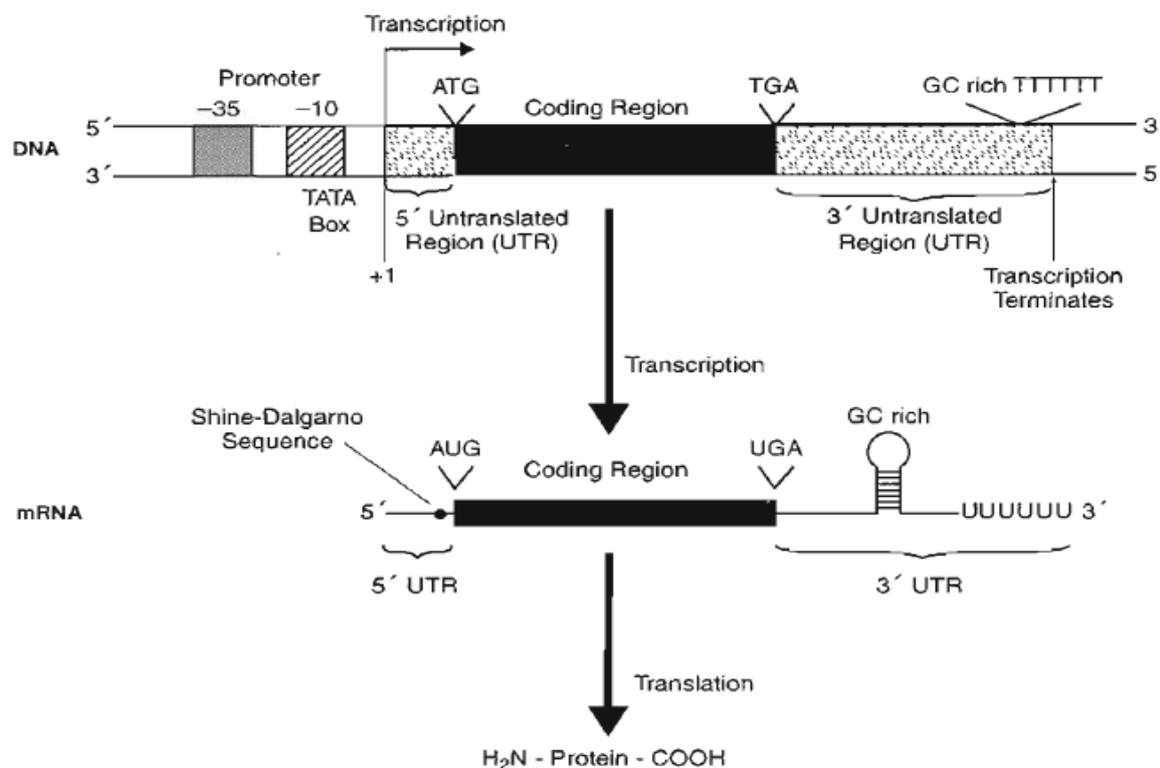
A- Rho-independent termination occurs when the newly formed RNA folds back on itself to form a GC-rich (strong bonds which make a mechanical stress) hairpin loop closely followed by **6-8 U** (weak bond) residues. These two structural features of the newly synthesized RNA promote dissociation of the RNA from DNA template.

B- Rho-dependent termination requires participation of rho factor (which is found at the end of the gene) (the RNA polymerase reads all of the DNA sequence of the gene while transcribing it which means even if it's in the middle it reads the sequence of the end of the gene), This protein binds to the newly formed RNA and moves toward the RNA polymerase that has paused at a termination site, Rho then displaces RNA polymerase from the 3' end of the RNA (breaking the hydrogen bonds).



5- Transcription and translation can occur simultaneously in bacteria because there is no processing of prokaryotic mRNA (generally no introns), ribosomes can begin translating the message even before transcription is complete.

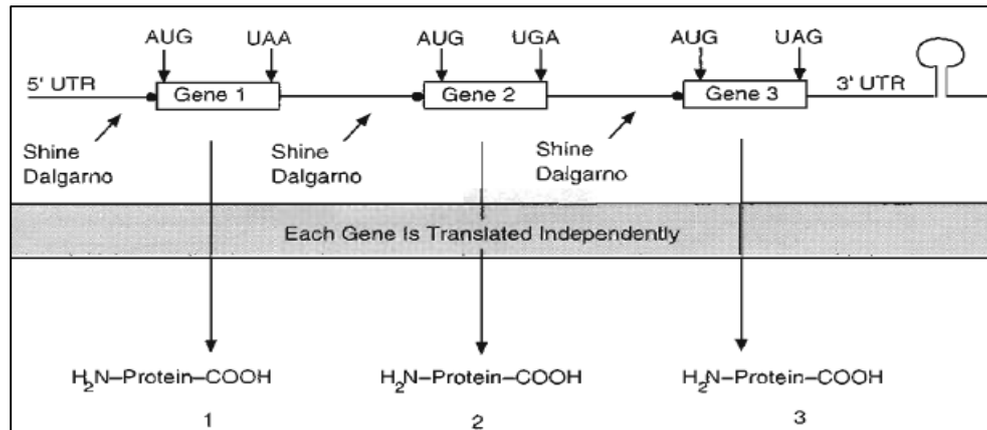
- ✓ Ribosomes (16s rRNA) bind to a sequence called Shine-Dalgarno sequence in the 5' untranslated region (UTR) of the mRNA.
 - ✓ Protein synthesis begins at an AUG codon at the beginning of the coding region and continues until the ribosome reaches a stop codon at the end of the coding region.
 - ✓ Either it's a eukaryotic or a prokaryotic cell the translation always begins at the genetic AUG codon (for the methionine amino acid)
- 6- The ribosome translates the message in the 5' to 3' direction, synthesizing the protein from amino terminus to carboxyl terminus.



A prokaryotic transcription unit

- ✓ The mRNA produced by the gene shown above is a **monocistronic** message (mRNA representing one gene). That is, it is transcribed from a single gene and codes for only a single protein.
- ✓ The word **cistron** is another name for a gene. Some bacterial operons produce **polycistronic** messages. In these cases, related genes grouped together in the DNA are transcribed as one unit.
- ✓ The mRNA in this case contains information from several genes and codes for several different proteins
- ✓ polycistronic messages are very very very very very rare for eukaryotic cells but some viruses can produce them.

Prokaryotic polycistronic message codes for several different proteins



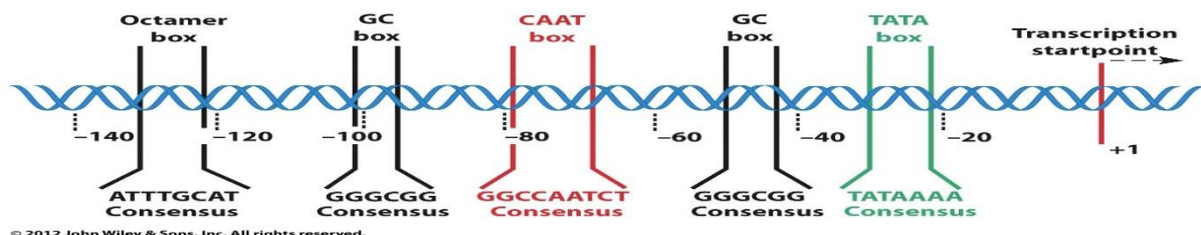
Production of eukaryotic mRNA:

- ✓ In eukaryotes, most genes are composed of coding segments (exons) interrupted by noncoding segments (introns).
- ✓ Both exons and introns are transcribed in the nucleus (because the RNA polymerase doesn't recognize them) so that pre-mRNA is formed
- ✓ Introns are removed during processing of the RNA molecule in the nucleus.
- ✓ In eukaryotes, all mRNA is monocistronic.
- ✓ The mature mRNA is translated in the cytoplasm.

Transcription of a typical eukaryotic gene occurs as follows:

1. With the help of proteins called transcription factors, RNA polymerase II recognizes and binds to the promoter region. The basal promoter region of eukaryotic genes usually has two consensus sequences called the **TATA box** (also called **Hogness box** / موقعه عند -25) and the **CAAT box**(between -70 to -90).
2. RNA polymerase II separates the strands of the DNA over a short region to initiate transcription and read the DNA sequence. The template strand is read in the 3' to 5' direction as the RNA product (the primary transcript) is synthesized in the 5' to 3' direction.

A Typical RNA Polymerase II Promoter



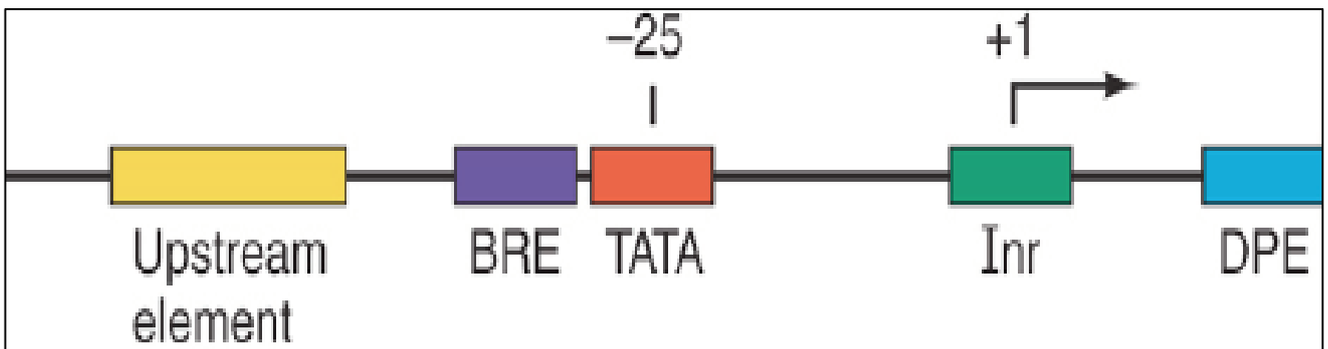
- **Class II promoters** (most similar to bacterial promoters)
 - ✓ Common type of promoter (most genes use this)
 - ✓ Many variations, but "consensus" has a "Core "+ "Upstream" Core (3 elements):
 1. "TATA box" (5'-TATA-3')(-25)
 2. TFIIB recognition element (BRE)
 3. Initiator box (Inr) with an "A" at +1, most common
 - ✓ Downstream promoter element (DPE, less common)
 - ✓ Core promoter is recognized by general TFs that associate with RNA pol to form a preinitiation complex at great majority of promoters
 - ✓ At least one of these elements is missing in most promoters

e.g., highly expressed specialized genes tend to have TATA boxes, but promoters for housekeeping genes tend to lack them because we don't need it to be highly active all time so they have caat box only and because of that it's difficult for RNA polymerase and transcription factors to recognize the initiation site so that the transcription of the housekeeping gene is a slow process and (منتج قليل ولكن في استمرارية يعني لوقت طويل) .

✓ Upstream elements: quite varied in number and can be **orientation-independent** (في عنا بوكسين فمش مهم مين الاول فيهم) (but relatively **position-dependent**) & recognized by other TFs (relatively **gene-specific**) that participate in initiation at smaller sub-sets of promoters.

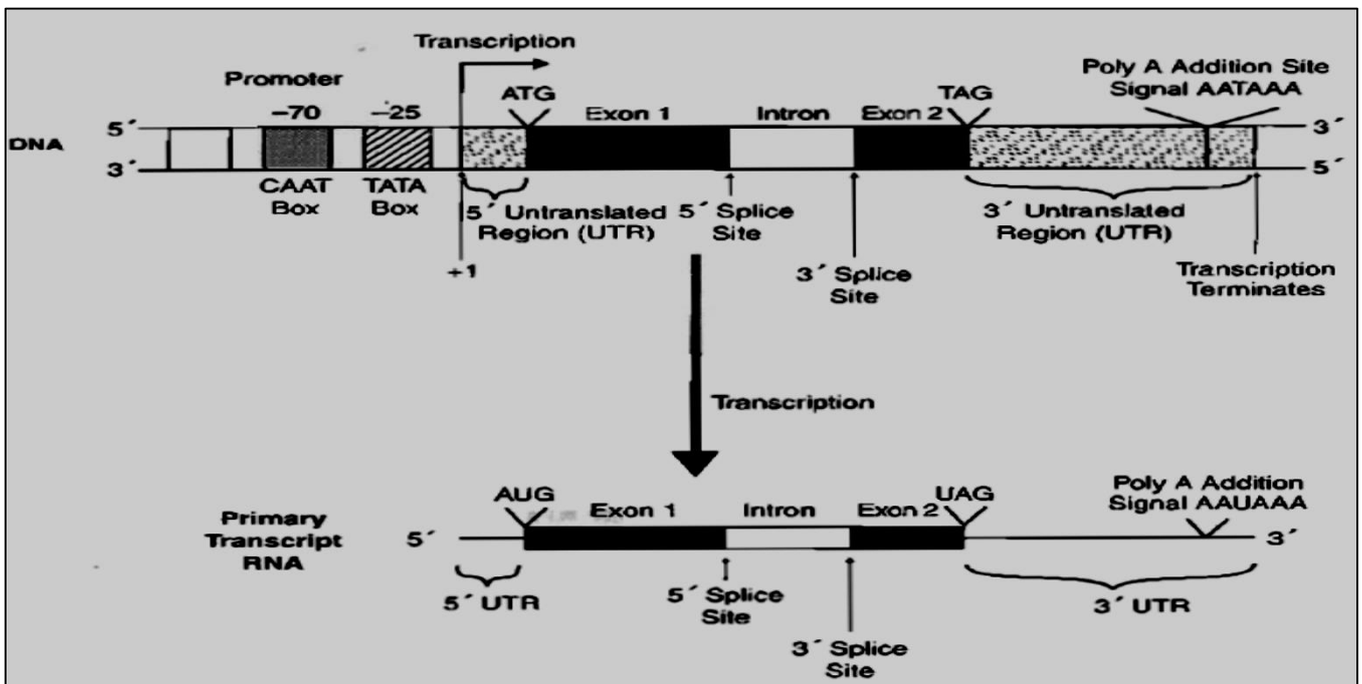
1. GC box (GC rich)
2. CAAT box (5'-CCAAT-3)

TFIIB recognition element; Inr: initiator box; DPE: downstream promoter element



3. RNA polymerase II ends transcription when it reaches a termination signal.
 - These signals are not well understood in eukaryotes.

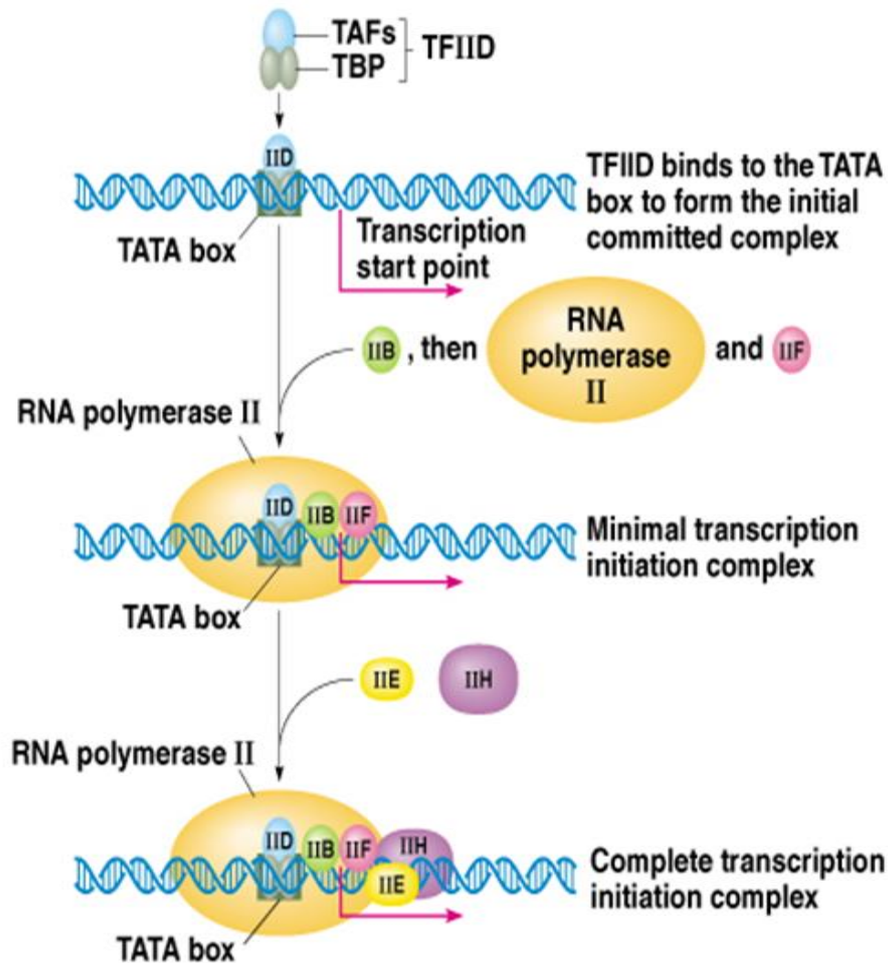
A eukaryotic transcription unit



- ❖ Exon is coding sequence and intron is non-coding sequence, and the exon are interrupted with intron as you see in the picture so that the introns must be removed by ribozymes to give continued exonic sequence.

Order of binding is: IID + IIA + IIB + RNA poly. II + IIF + IIE + IIH

a) Assembly of preinitiation complex



- 1- TBP (TATA binding protein) in TFIID (transcription factor II D) binds to the TATA box to form the initial complex
 - 2- TFIIA and TFIIB are recruited with TFIIB binding to the BRE
 - 3- RNA Pol II-TFIIF complex is then recruited
 - 4- TFIIE and TFIIH then bind upstream of Pol II to form the pre-initiation complex
- Promoter melting using energy from ATP hydrolysis by TFIIH)
 - Promoter escapes after the phosphorylation of the C-terminal domain tail

❖ الجدول الي جاي مهم ☺

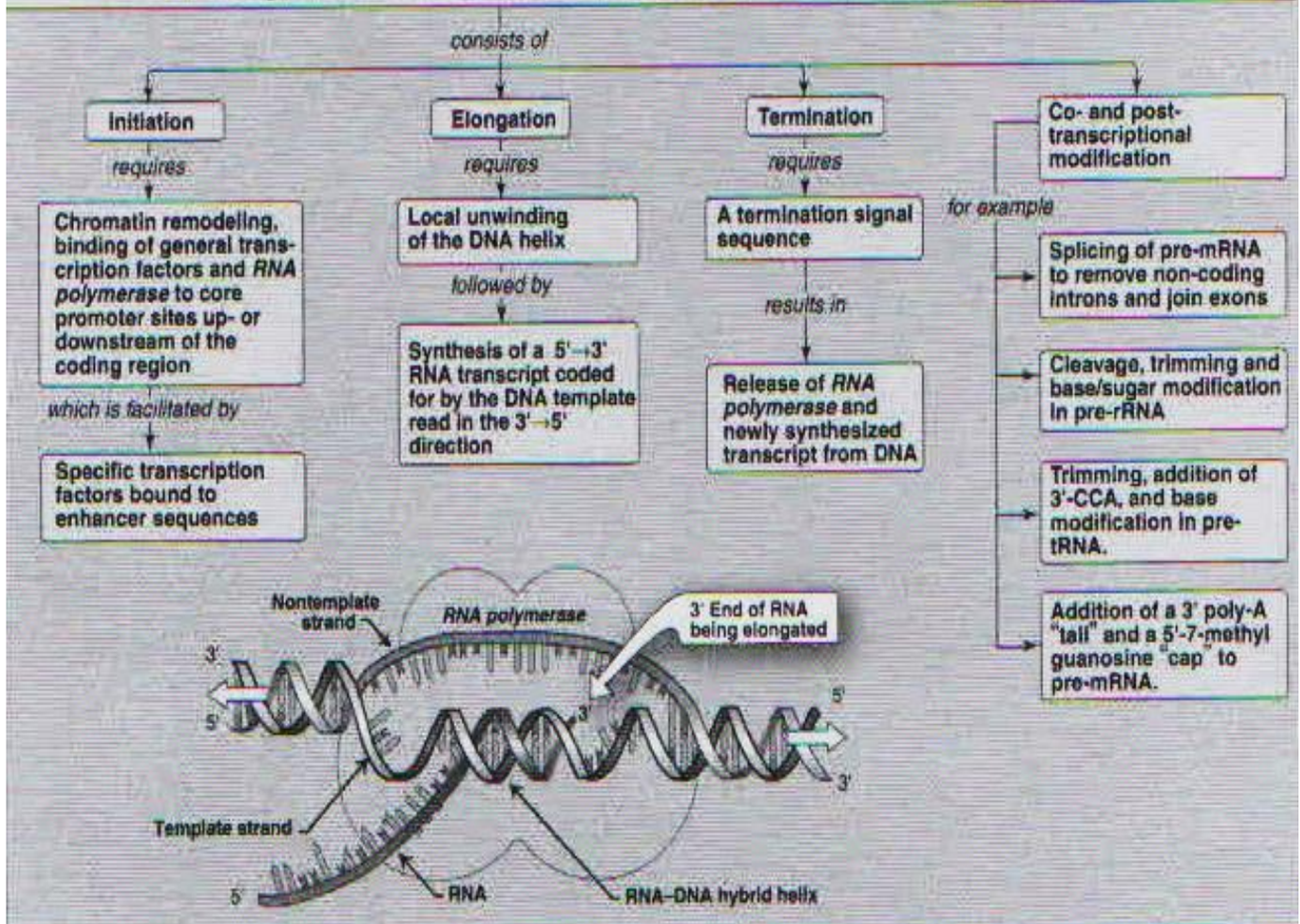
- ❖ TFIID-TAF It's considers a positive and negative regulation so if it recognizes non-TATA element the rate of transcription increases (والعكس صحيح)
- ❖ TFIIE recruits (يستدعي) TFIIH and stimulates its activities
- ❖ TFIIH work as helicase ATPase and kinase +promoter melting (destroying hydrogen bonds) + promoter clearance by phosphorylation of carboxylic terminal domain
- ❖ ال RNA polymerase بعد ما يبدأ تصنيع ال mRNA يشكك في نفسه انه بدأ من المكان الصحيح ام لا ويعمل ايقاف للعملية (abortion) ويبرجع بشيك على نقطة البداية ويعمل هالاشي اكثر من مره

Table 11.3

General Transcription Initiation Factors

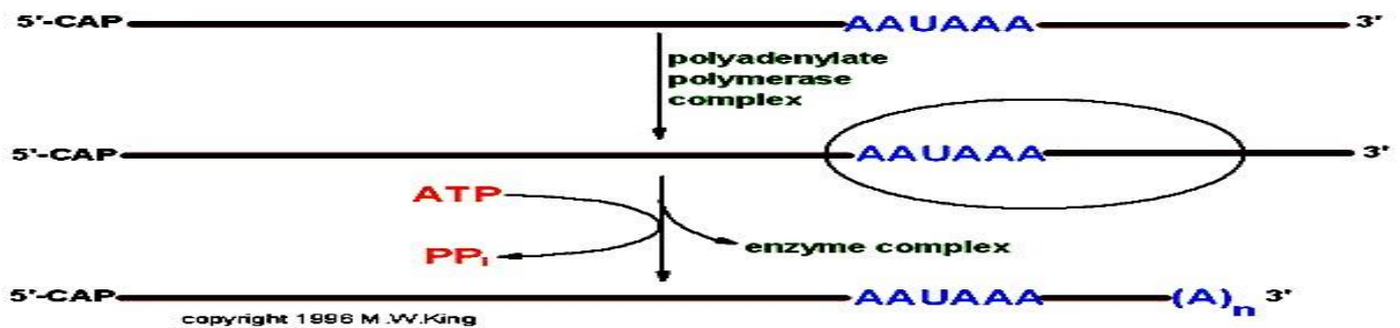
Factor	Subunits	Size (kDa)	Function
TFIID-TBP	1	27	TATA box recognition, positioning of TATA box DNA around TFIIB and Pol II
TFIID-TAF _{II} s	14	15–250	Core promoter recognition (non-TATA elements), positive and negative regulation
TFIIA	3	12, 19, 35	Stabilization of TBP binding; stabilization of TAF–DNA binding
TFIIB	1	38	Recruitment of Pol II and TFIIF; start-site recognition for Pol II
TFIIF	3	156 total	Promoter targeting of Pol II
TFIIE	2	92 total	TFIIH recruitment; modulation of TFIIH helicase ATPase, and kinase activities; promoter melting
TFIIH	9	525 total	Promoter melting; promoter clearance via phosphorylation of CTD

Eukaryotic Transcription: DNA-Directed RNA Synthesis



- ✓ A stretch of 20–250 residues is then added to the 3' end by the **polyadenylate polymerase** activity.

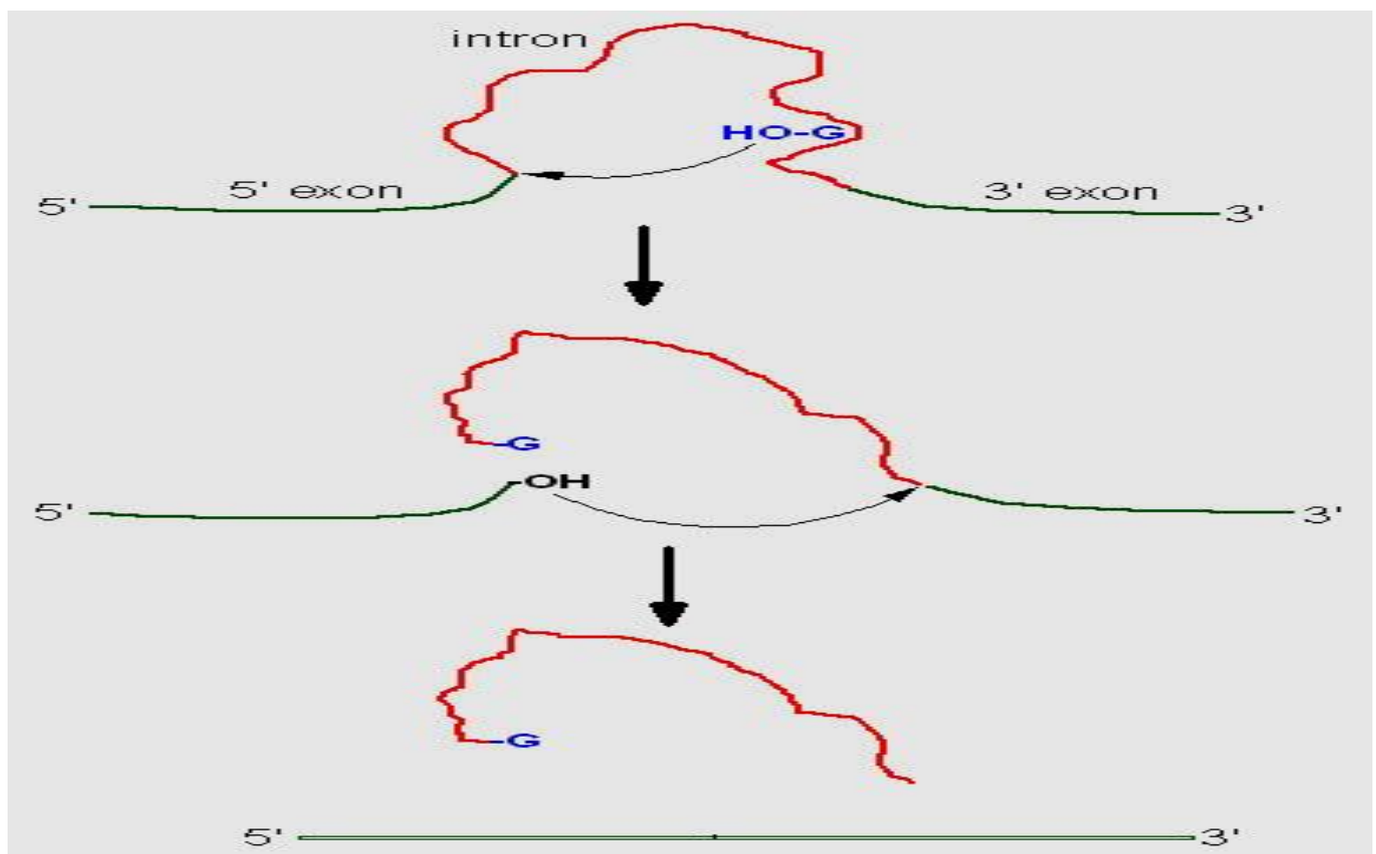
Polyadenylation of mRNAs



✚ Splicing of RNAs:

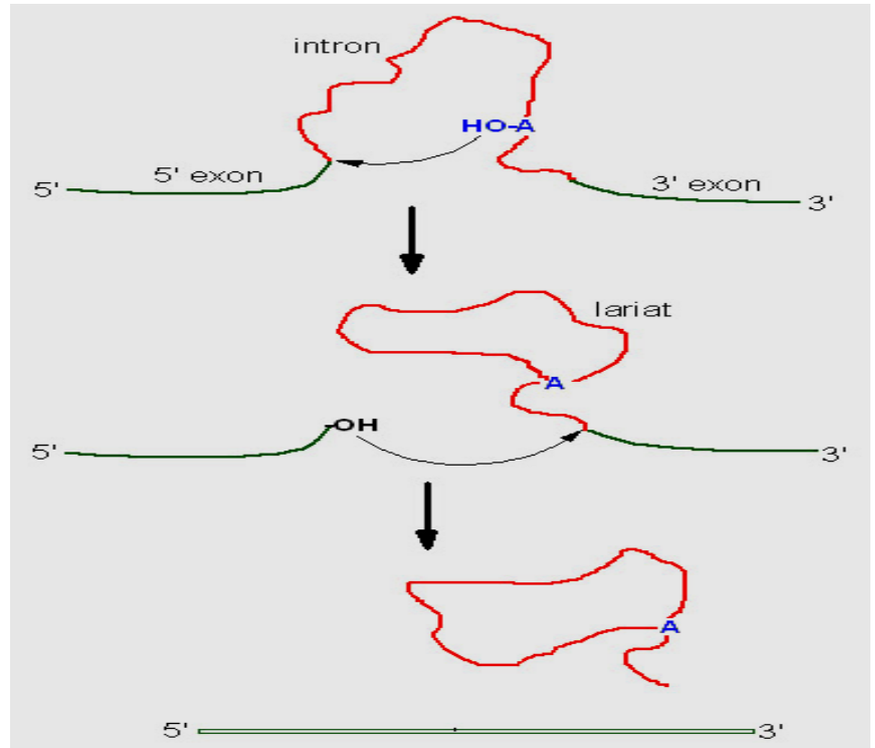
- ✓ There are several different classes of reactions involved in intron removal.
- ✓ The 2 most common are the group I and group II introns.
- ✓ Group I introns are found in nuclear, mitochondrial and chloroplast rRNA genes, group II in mitochondrial and chloroplast mRNA genes.
- ✓ Many of the group I and group II introns are self-splicing.
- ✓ Group I introns require an external guanosine as a cofactor.
- ✓ The 3'-OH of the guanosine nucleotide acts as a nucleophile to attack the 5'-phosphate of the 5' nucleotide of the intron.
- ✓ The resultant 3'-OH at the 3' end of the 5' exon then attacks the 5' nucleotide of the 3' exon releasing the intron and covalently attaching the two exons together.
- ✓ The 3' end of the 5' exon is termed the splice donor site and the 5' end of the 3' exon is termed the splice acceptor site.

Splicing by group 1 introns



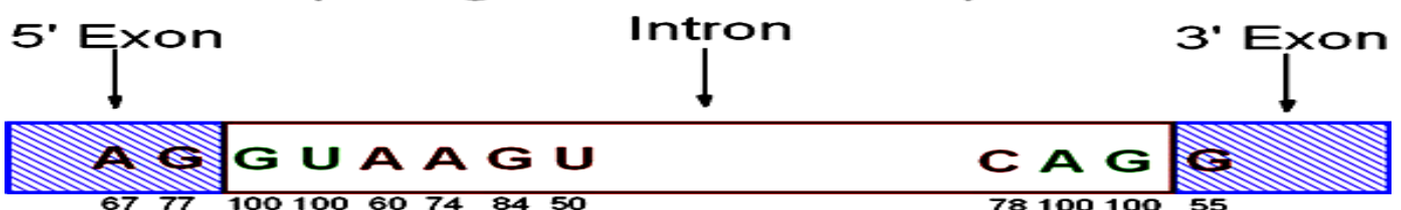
- ✓ Group II introns are spliced similarly except that instead of an external nucleophile, the 2'-OH of an adenine residue within the intron is the nucleophile.
- ✓ This residue attacks the 3' nucleotide of the 5' exon forming an internal loop called a lariat structure.
- ✓ The 3' end of the 5' exon then attacks the 5' end of the 3' exon as in group I splicing releasing the intron and covalently attaching the two exons together.

Splicing by group 2 introns



- ✓ The third class of introns is also the largest class found in nuclear mRNAs, that undergoes a splicing reaction similar to group II introns in that an internal lariat structure is formed.
- ✓ However, the splicing is catalyzed by specialized RNA– protein complexes called small nuclear ribonucleoprotein particles (snRNPs).
- ✓ The RNAs found in snRNPs are identified as U1, U2, U4, U5 and U6.
- ✓ Analysis of a large number of mRNA genes has led to the identification of highly conserved consensus sequences at the 5' and 3' ends of essentially all mRNA introns.
- ✓ The U1 RNA has sequences that are complimentary to sequences near the 5' end of the intron, its binding allows distinguishing the GU at the 5' end of the intron from other randomly placed GU sequences in mRNAs.
- ✓ The U2 RNA also recognizes sequences in the intron, in this case near the 3' end.
- ✓ The addition of U4, U5 and U6 RNAs forms a complex identified as the spliceosome (snRNA plus ~40 proteins) that removes the intron and joins the two exons together.
- ✓ U7 is involved in the production of **the correct 3' ends of histone mRNA which lacks poly (A) tail.**
- ✓ An additional mechanism of intron removal is the process of tRNA splicing.

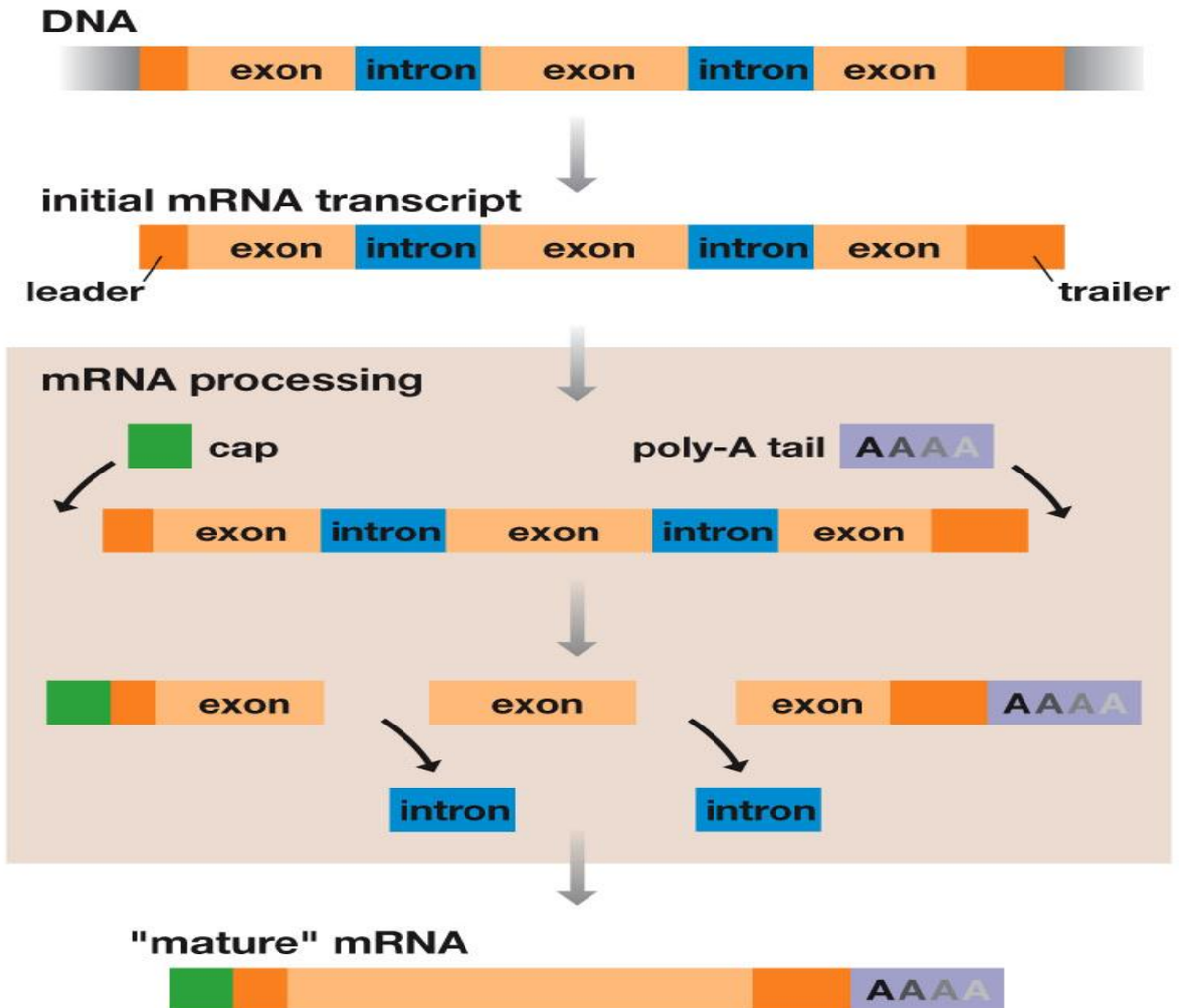
Splicing Consensus Sequences



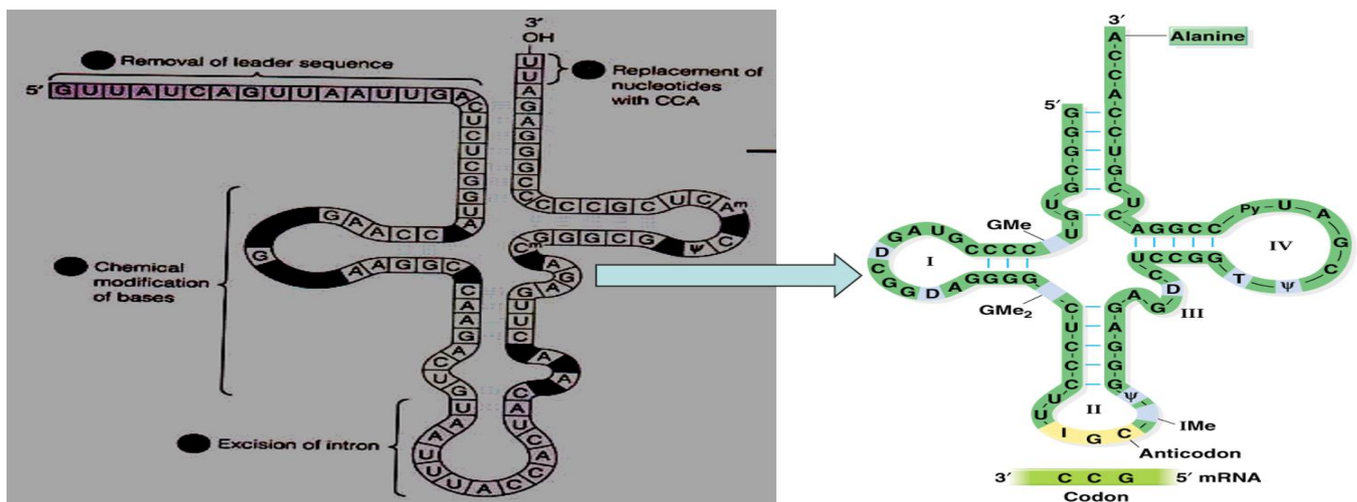
- ✓ These introns are spliced by a specific splicing endonuclease that involves a cut-and-paste mechanism.
- ✓ In order for tRNA intron removal to occur the tRNA must first be properly folded into its characteristic cloverleaf shape.
- ✓ Misfolded precursor tRNAs are not processed which allows the splicing reaction to serve as a control step in the generation of mature tRNAs.

RNA processing: pre-mRNA → mRNA

✚ Modifications of tRNA



- 1- Removal of 5' extrasequence
- 2- Addition of: - CCA at 3' end - Anticodon loop
- 3- Methylation of some bases



✚ rRNA is used to construct ribosomes

- ✓ Eukaryotic ribosomal RNA is transcribed in the nucleolus by RNA polymerase I as a single piece of 45S RNA, which is subsequently cleaved to yield 28S rRNA, 18S rRNA, and 5.8S rRNA.
- ✓ RNA polymerase III transcribes the 5S rRNA unit from a separate gene. The ribosomal subunits assemble in the nucleolus as the rRNA pieces combine with ribosomal proteins.
- ✓ Eukaryotic ribosomal subunits are 60S and 40S. They join during protein synthesis to form the whole 80S ribosome.

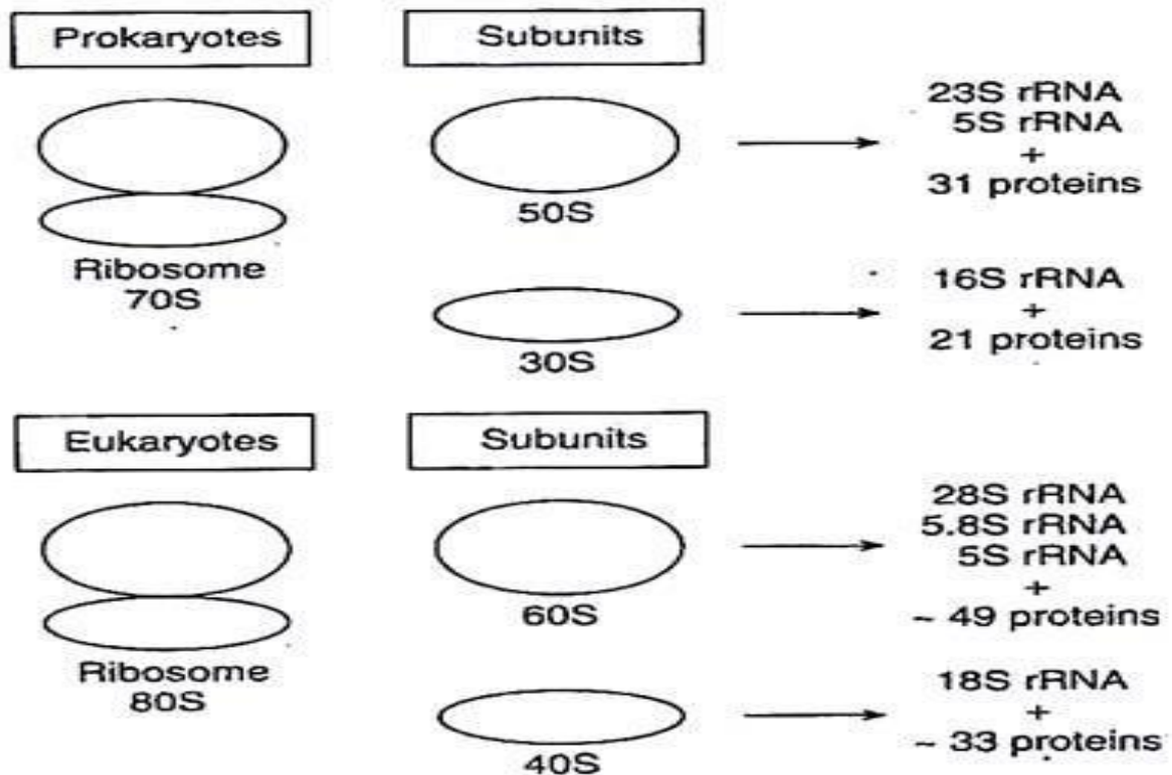


Fig. 2.47: Composition of typical prokaryotic and eukaryotic ribosomes