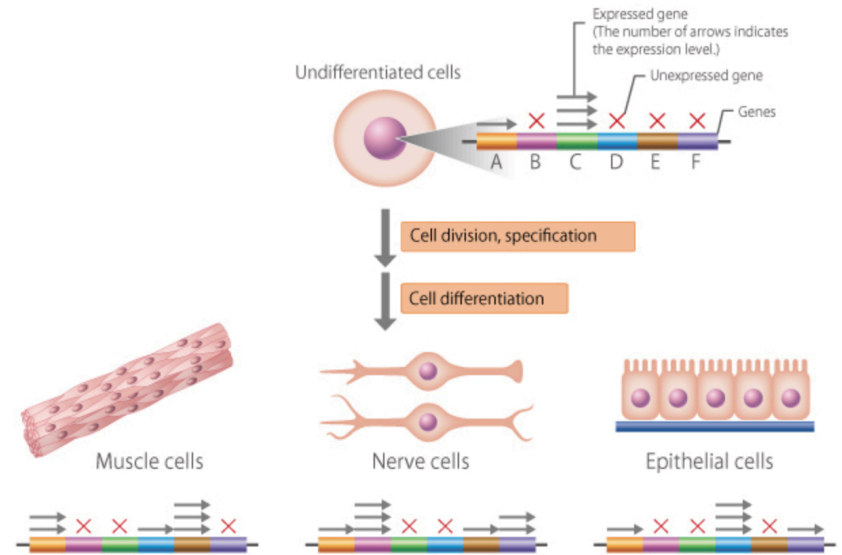
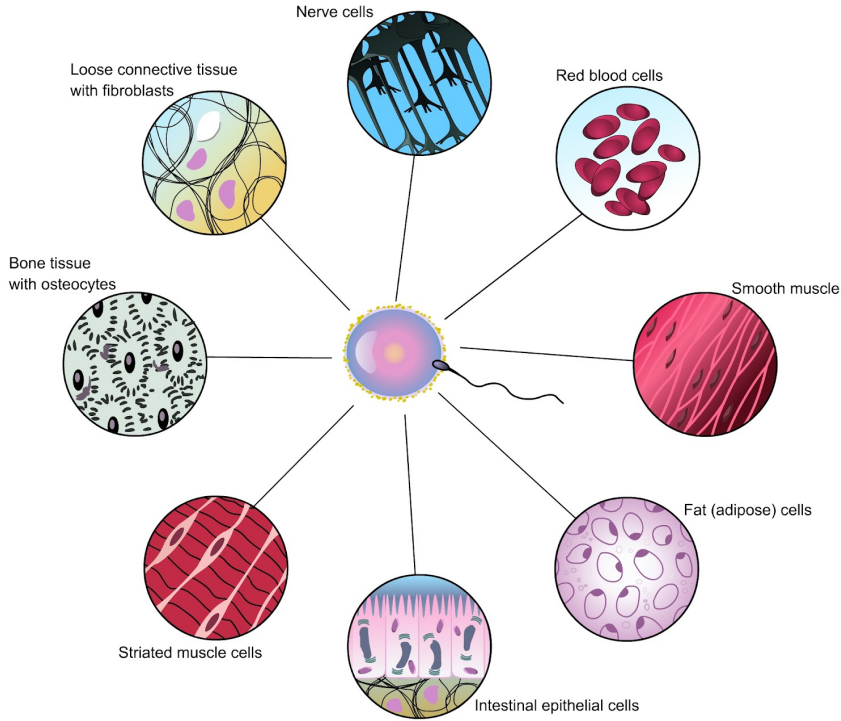


# **Principles of Gene Expression**

# I. Introduction

Regulation of gene expression is intrinsically linked to multicellularity and cell-specification in higher eucaryotes



## II. Principles of gene regulation

### Principle #1 : constitutive vs regulated gene expression

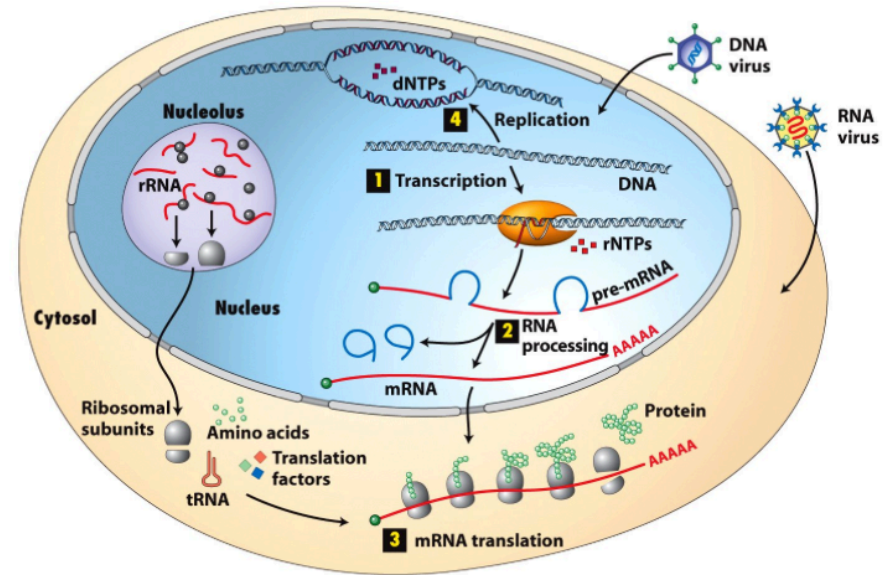
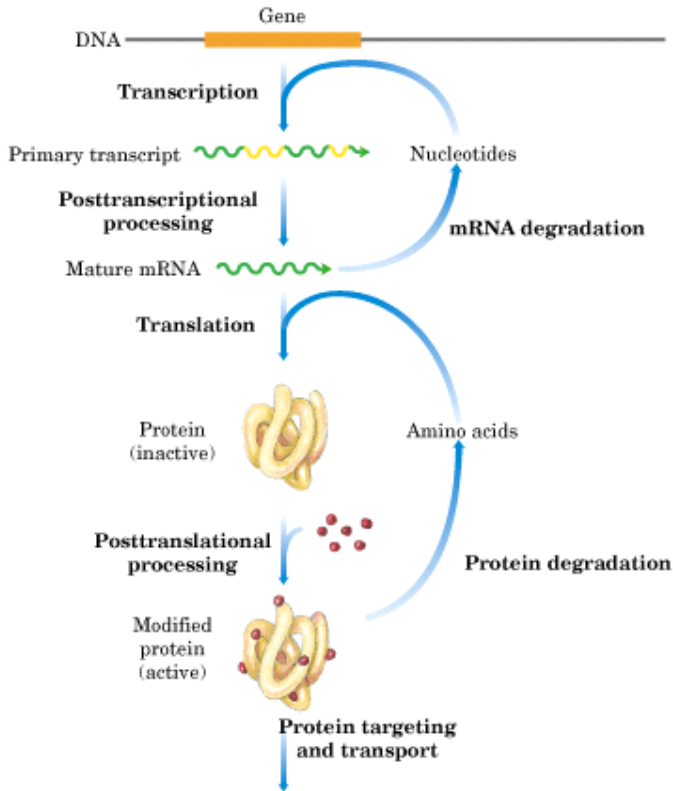
There are approx 10.000 genes expressed in an eukaryotic cell  
(4.000 in yeast)

Most of the genes code for polypeptides that are required at all times at a more or less constant level, expressed at low levels, common to different cell types (*housekeeping genes*)

A minority of mARNs are expressed at high levels, they are specific to the cell type or conditions (*luxury genes*)

## II. Principles of gene regulation

**Principle #2 :** Gene regulation can be achieved at several levels along polypeptide synthesis (different in prokaryotes and eukaryotes).





## II. Principles of gene regulation

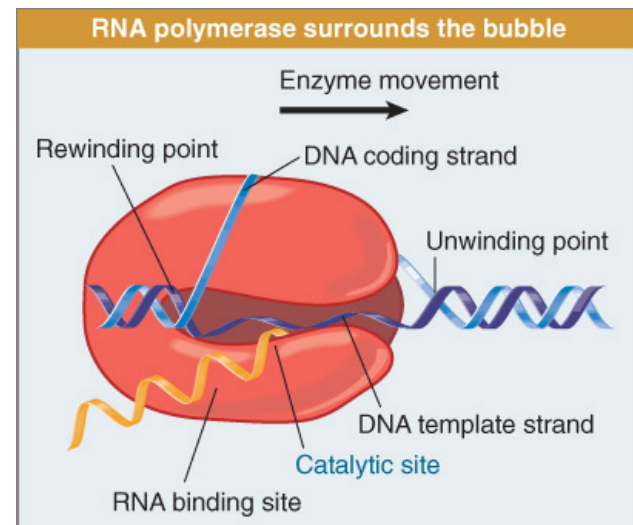
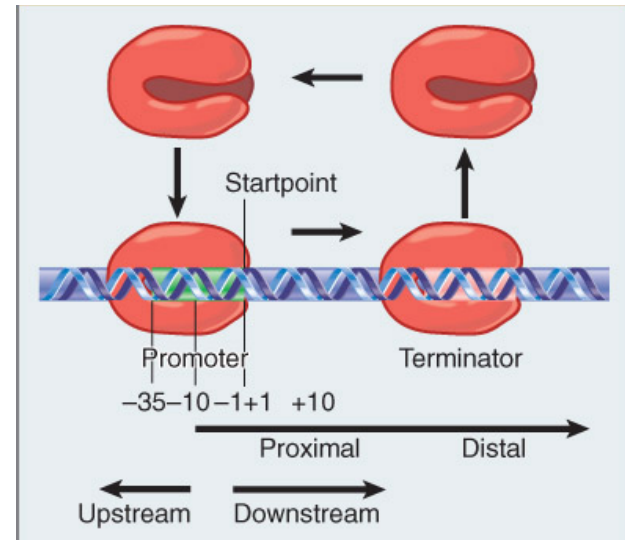
**Principle #3** : Transcription proceeds through a series of defined processes : initiation, elongation and termination

RNA polymerase recognizes **promoter sequences (initiation)**.

RNA polymerase synthesizes RNA complementary to the template DNA (**elongation**)

RNA synthesis proceeds through terminator sequences (**termination**)

The promoter, the transcribed region and the terminator define a **transcription unit**



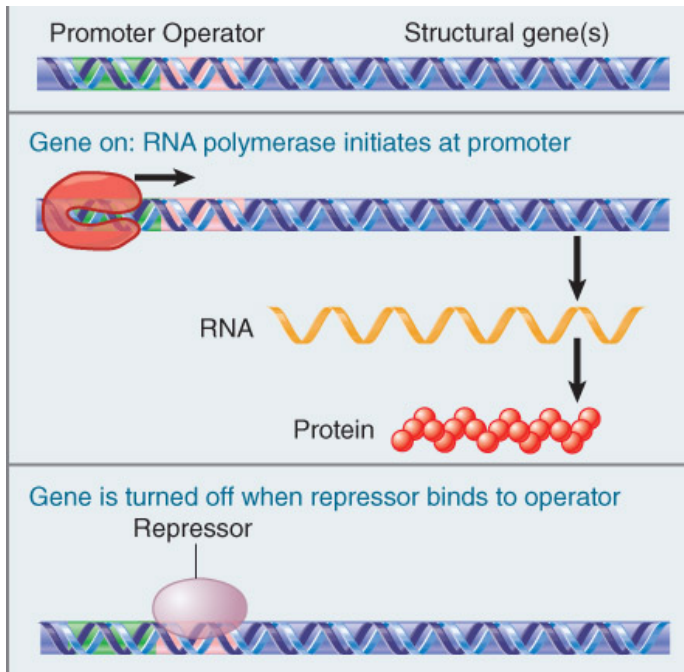
## II. Principles of gene regulation

**Principle #4** : Transcription initiation is regulated **by trans-acting proteins** (activators or repressors) that bind **to cis-acting sites** in or near promoters

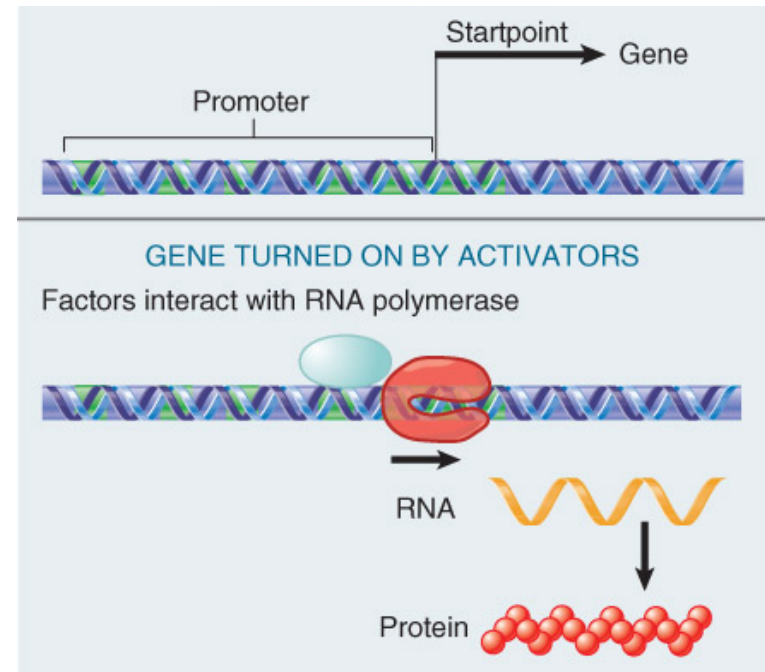
**Repressors** bind to **operators** in DNA to prevent or reduces the RNA polymerase activity : (*i.e.*, **negative regulation**)

**Activator** binds to DNA sites called **enhancer** to enhance the RNA polymerase activity. (*i.e.*, **positive regulation**)

### Negative regulation

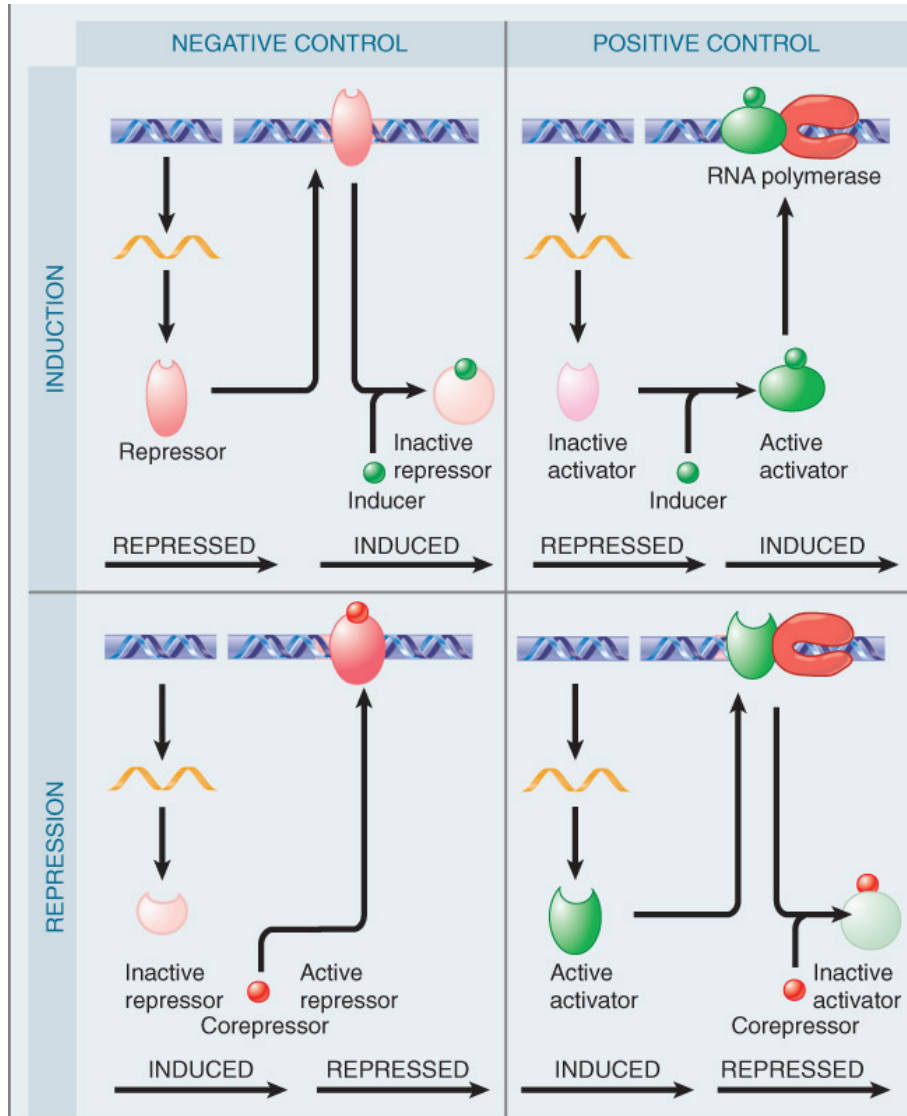


### Positive regulation



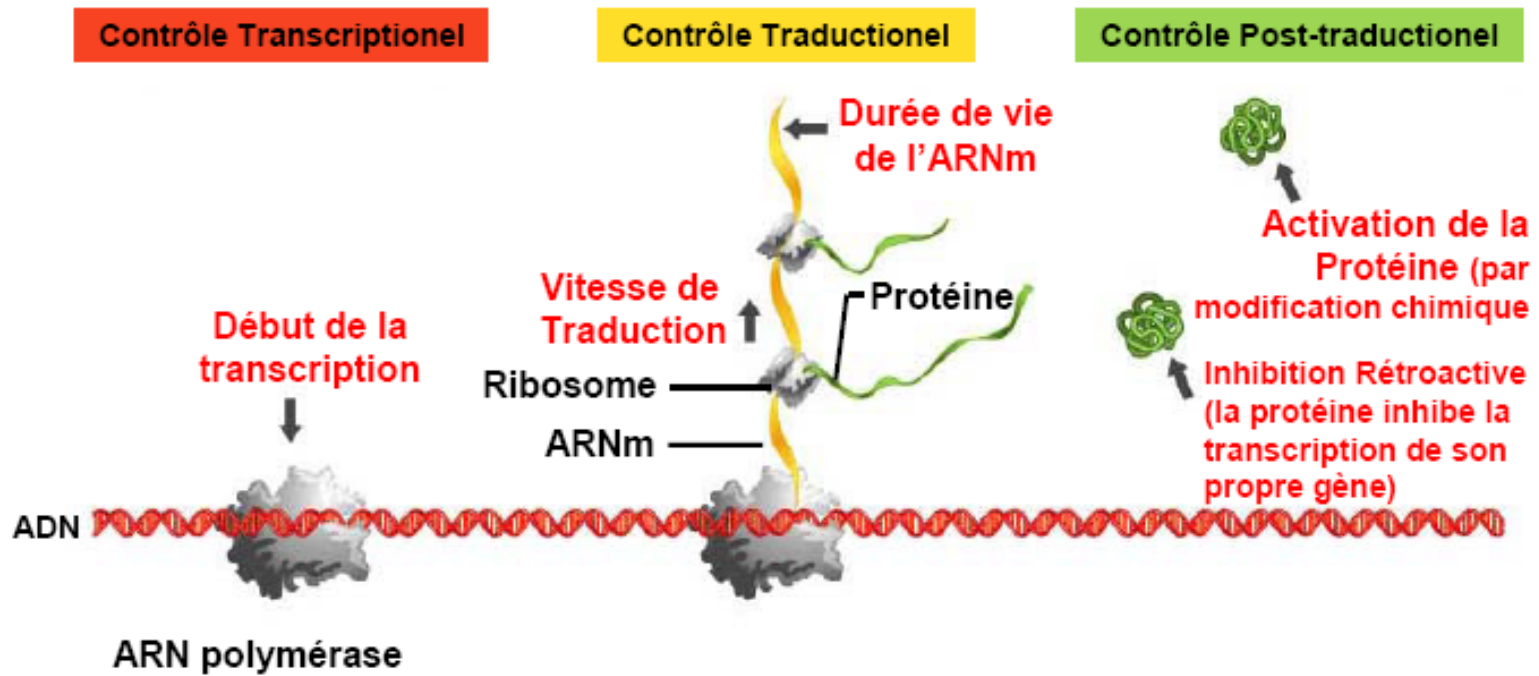
## II. Principles of gene regulation

**Principle #5 :** Activation or repression by activators or repressors can be regulated by **co-regulators** (small molecules in procaryotes)



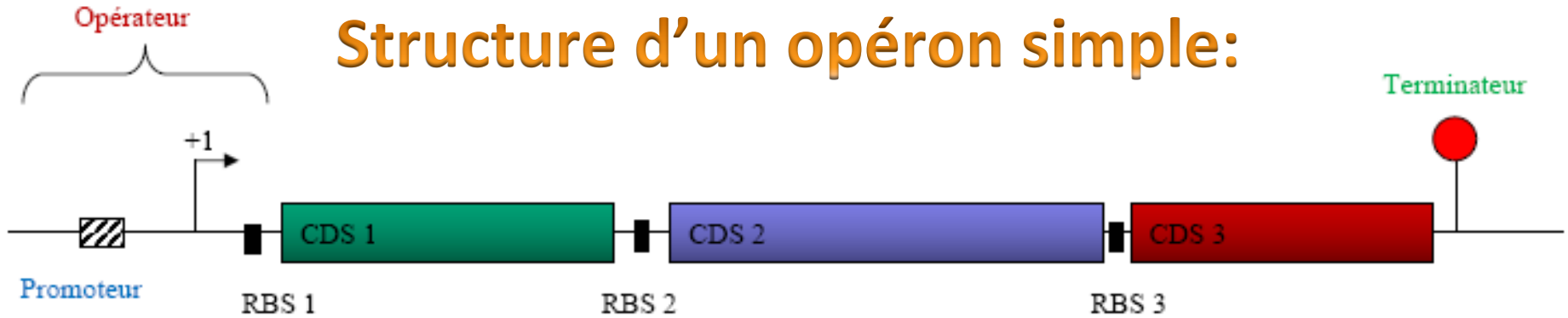
### III. Gene Regulation in Prokaryotes

**Principle** : Gene regulation in Prokaryotes is achieved at the **transcription initiation level**



### III. Gene Regulation in Prokaryotes

Prokaryotic genes are expressed and regulated as **operons**



**Operator** : DNA sequences (binding sites) involved in transcription initiation control

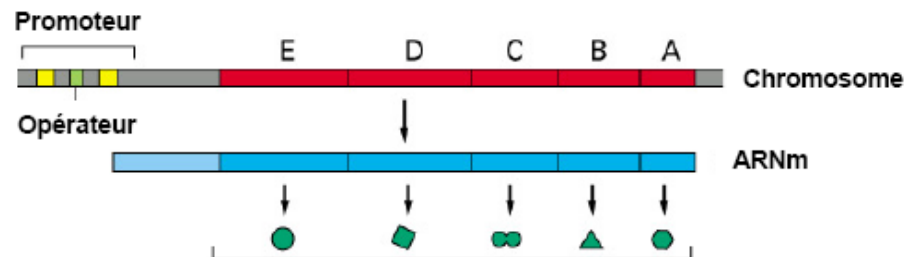
Promoter : binding of RNA polymerase

+1 : transcription start

RBS : ribosome binding site

CDS : coding sequence

**Terminator** : ends transcription



### III. Gene Regulation in Prokaryotes

Bacterial promoters are defined by 4 core elements

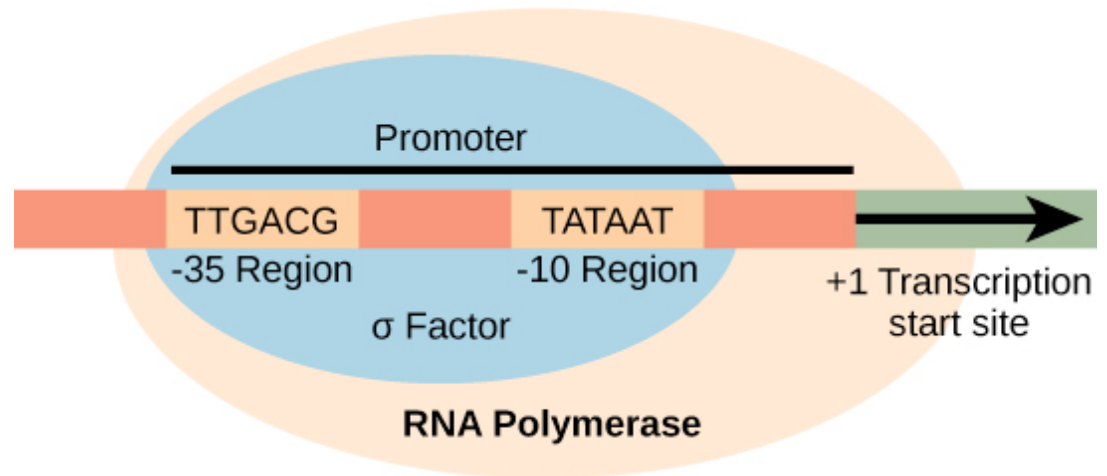
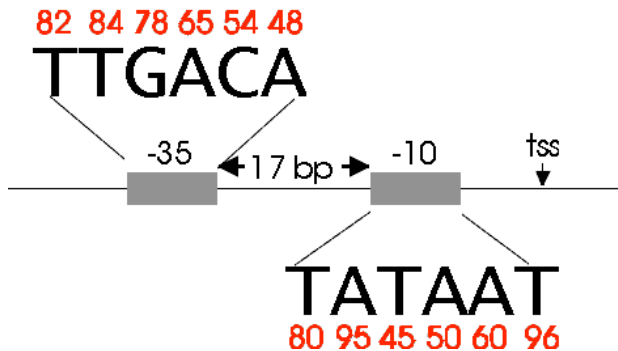
TSS : transcription start site (purine in >90%)

TATA box at around -10 from the TSS

-35 Box

An insert region between the boxes of 16-18 nucleotide

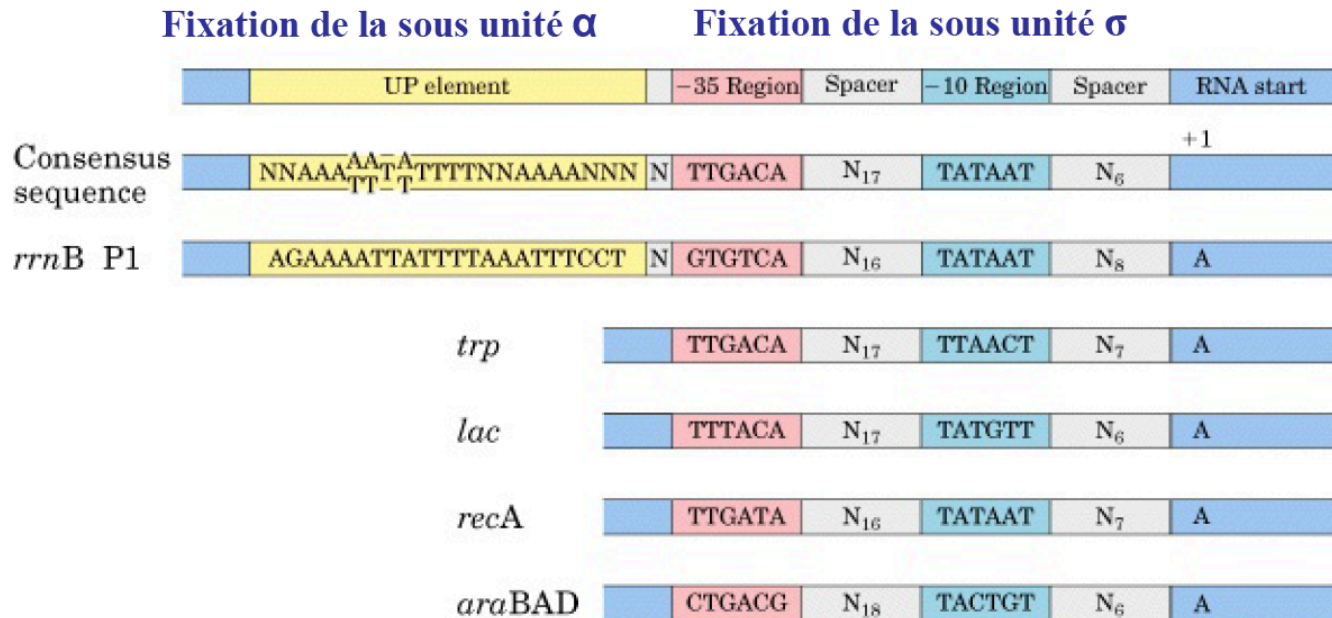
Typical Bacterial Promoter



### III. Gene Regulation in Prokaryotes

Divergence from canonical sequences and structure defines how efficiently a promoter will initiate transcription (promoter strength : weak or strong).

Mutations in promoter regions can affect promoter strength (mutations up/down)



### III. Gene Regulation in Prokaryotes

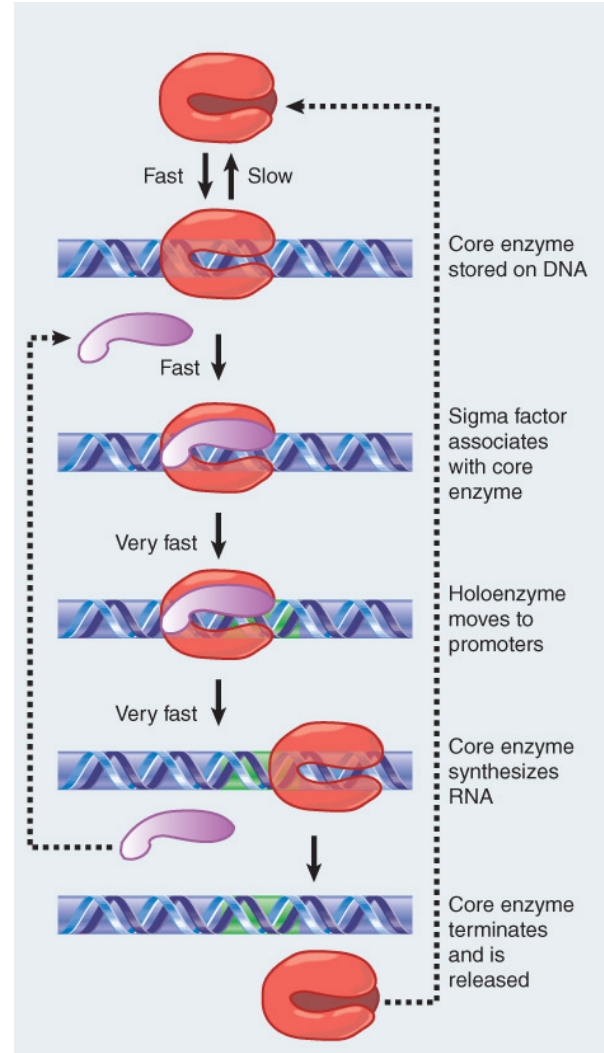
RNA polymerase requires  $\sigma$ -factors for efficient binding to DNA-sequences at TSS

The  $\sigma$ -factor has specific domains that recognize canonical elements in the promoter

The  $\sigma$ -factor controls the affinity of the polymerase for the promoter sequences

Recruitment of RNA polymerases to TSS is regulated by activating and repressing trans-acting factors

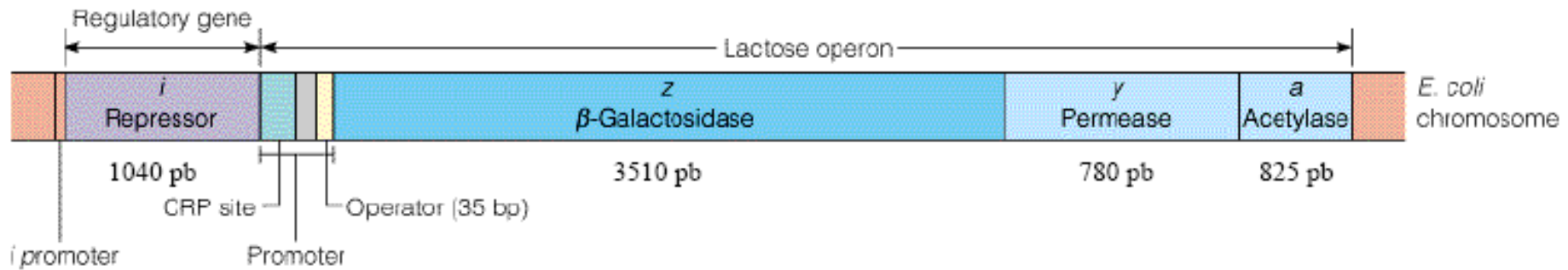
Most cis-acting elements are close to the TSS





### III. Gene Regulation in Prokaryotes

**Lactose operon** encodes enzymes involved in  **$\beta$ -galactosides catabolism**



LacZ :  $\beta$ -galactosidase (hydrolyzes  $\beta$ -galactoside)

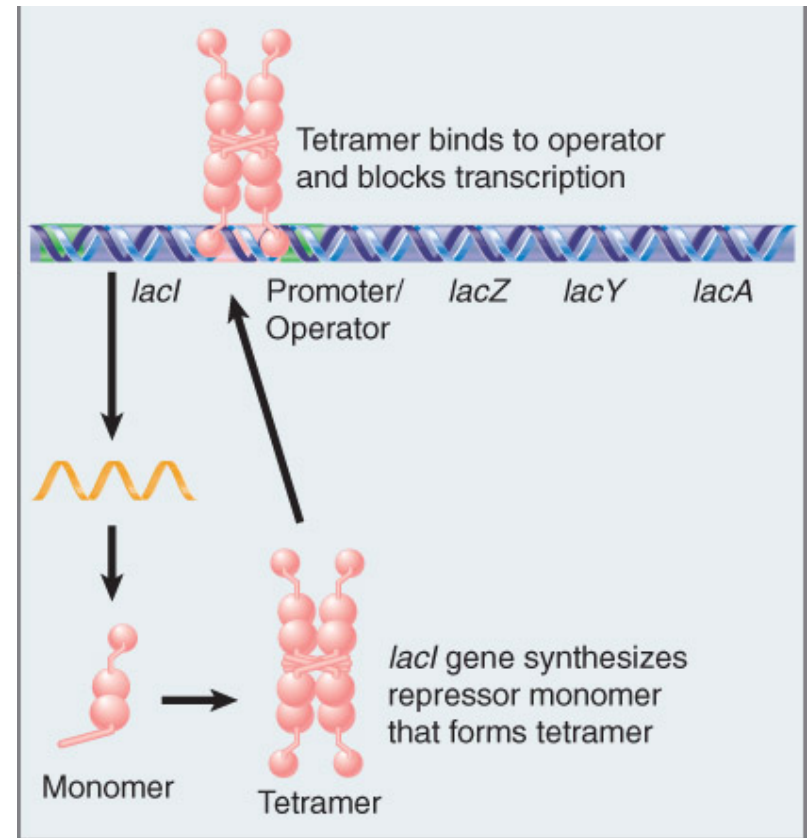
LacY : Permease (imports  $\beta$ -galactosides inside *E. coli*)

LacA : trans-Acetylase ( acetylates  $\beta$ -galactosides)

### III. Gene Regulation in Prokaryotes

The *lac repressor* is transcribed from its own promoter Pi

**LacI** binds the Plac (operator) and prevents transcription initiation



### III. Gene Regulation in Prokaryotes

*LacI* is itself inactivated by an inducer (allolactose)

#### Lactose paradox

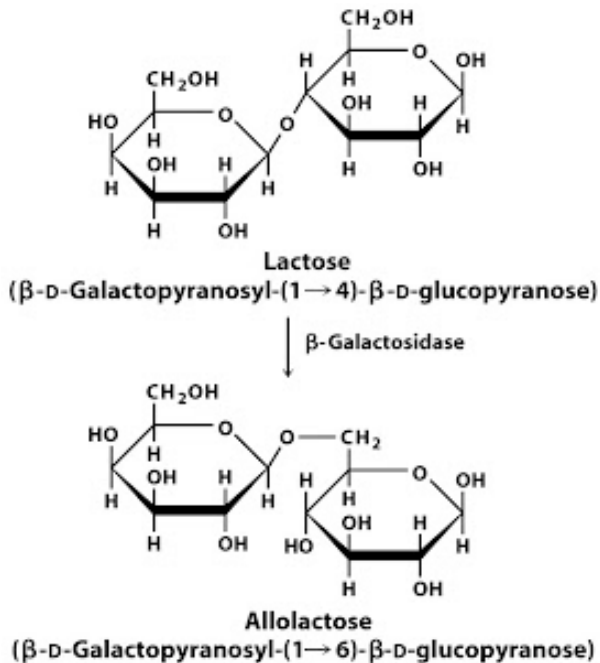
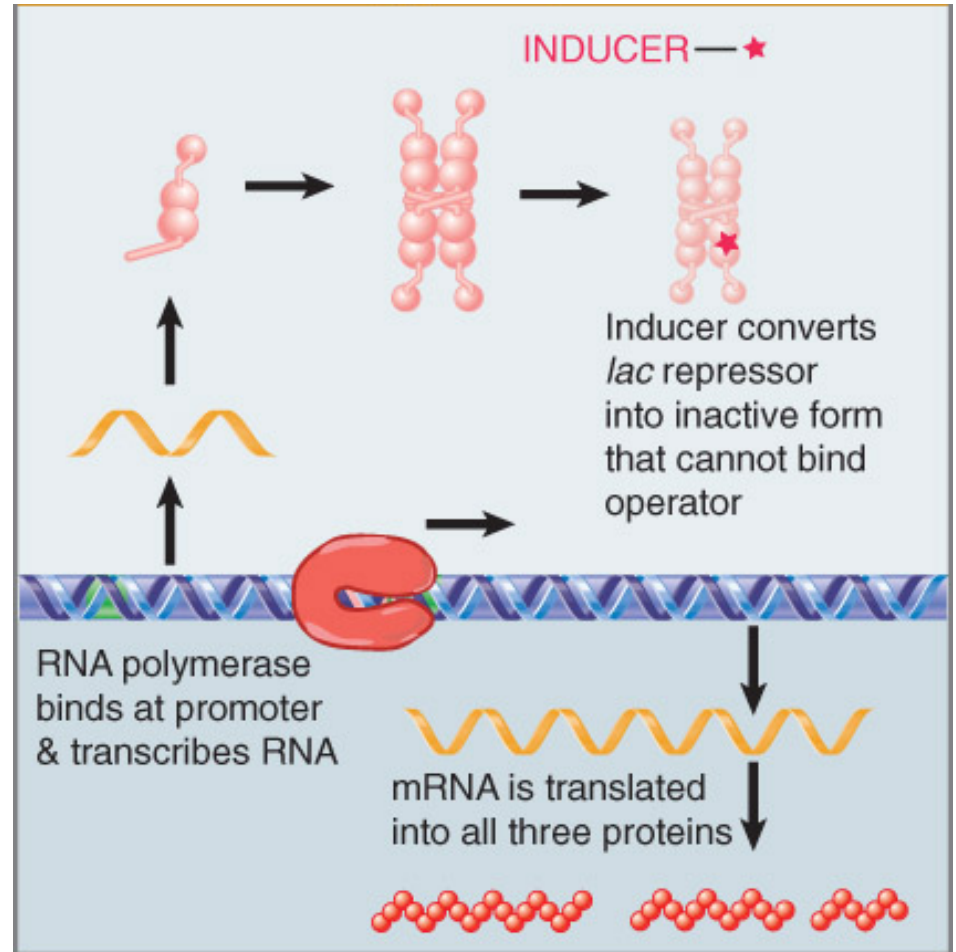
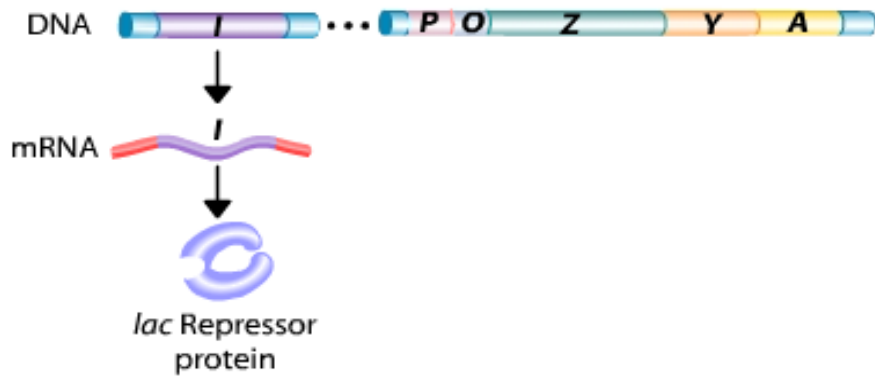


Figure 21-18 Principles of Biochemistry, 4/e  
© 2005 Pearson Prentice Hall, Inc.

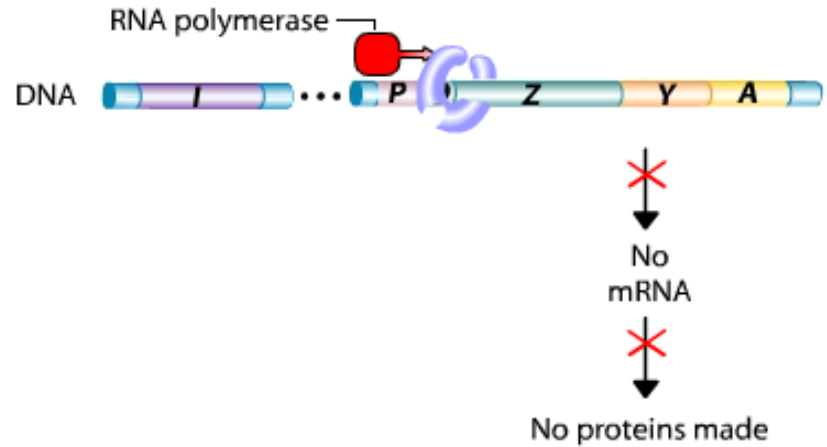


### III. Gene Regulation in Prokaryotes

## Glucose/ No Lactose



The Lac repressor protein, encoded by the *I* gene, is expressed in the absence or presence of lactose.



In the absence of lactose, the Lac repressor binds to the *lac* operator site.

Since RNA polymerase is unable to transcribe the *lac* structural genes, the corresponding proteins are not made.

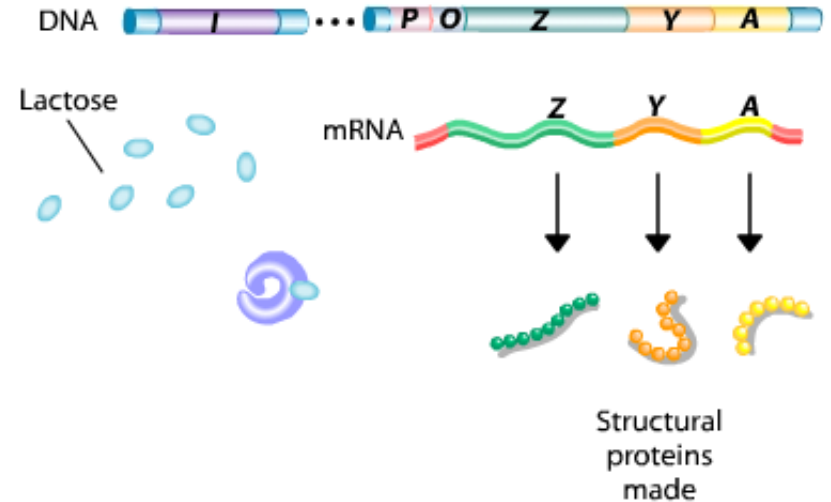
### III. Gene Regulation in Prokaryotes

## Lactose



When lactose is present in the cell medium, it binds to the allosteric site of the Lac repressor. This changes the conformation of the repressor.

In this conformation, the repressor can no longer bind to the *lac* operator site.

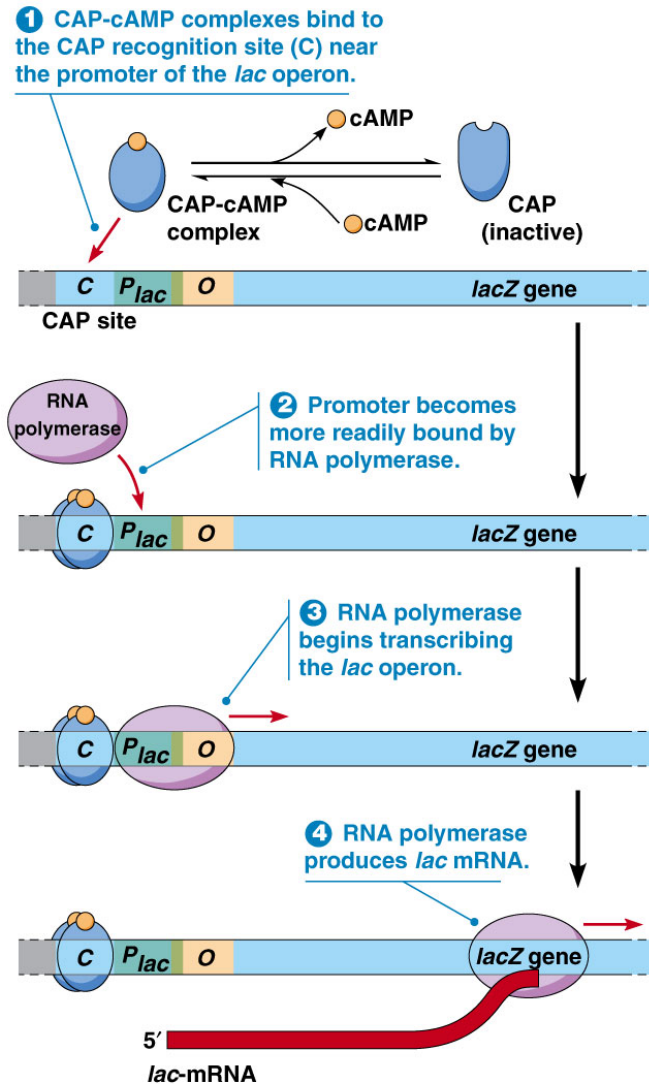
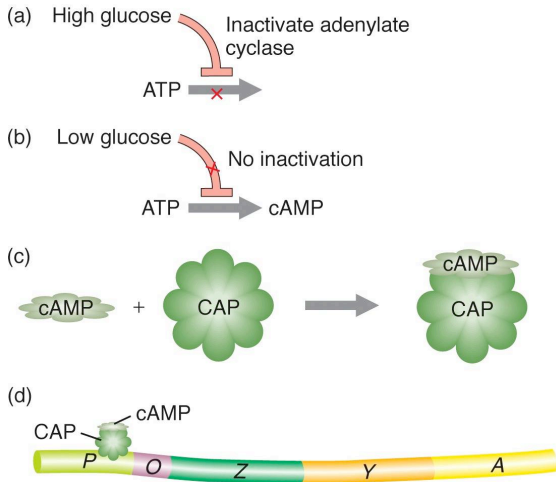


Without the repressor blocking its way, RNA polymerase is able to transcribe the structural genes.

Thus, in the presence of lactose, the *lac* structural genes are expressed. The proteins encoded by the Z and Y genes are required for the metabolism of lactose.

### III. Gene Regulation in Prokaryotes

Preferential use of Glucose over lactose is achieved through the **cAMP Receptor Complex (CRP)**



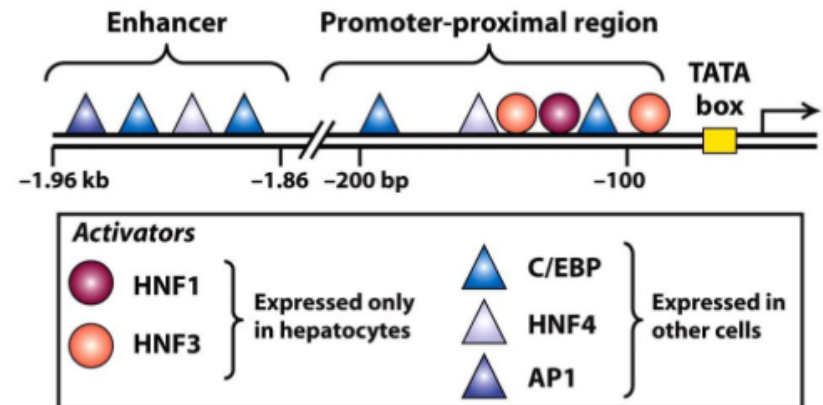
## IV. Gene Regulation in Eukaryotes

Unlike bacterial cells and most single cell eukaryotes, cells in multicellular organisms have relatively few genes that are directly and reversibly regulated by environmental conditions

Instead multicellular organisms have many different, specialized cells. Hence, **tissue-specific gene control** is important for development and differentiation.

every gene has more than one gene regulator (all of which must be on for the gene to function).

Eukaryotic regulatory elements are usually several Kb away from the promoter



# IV. Gene Regulation in Eukaryotes

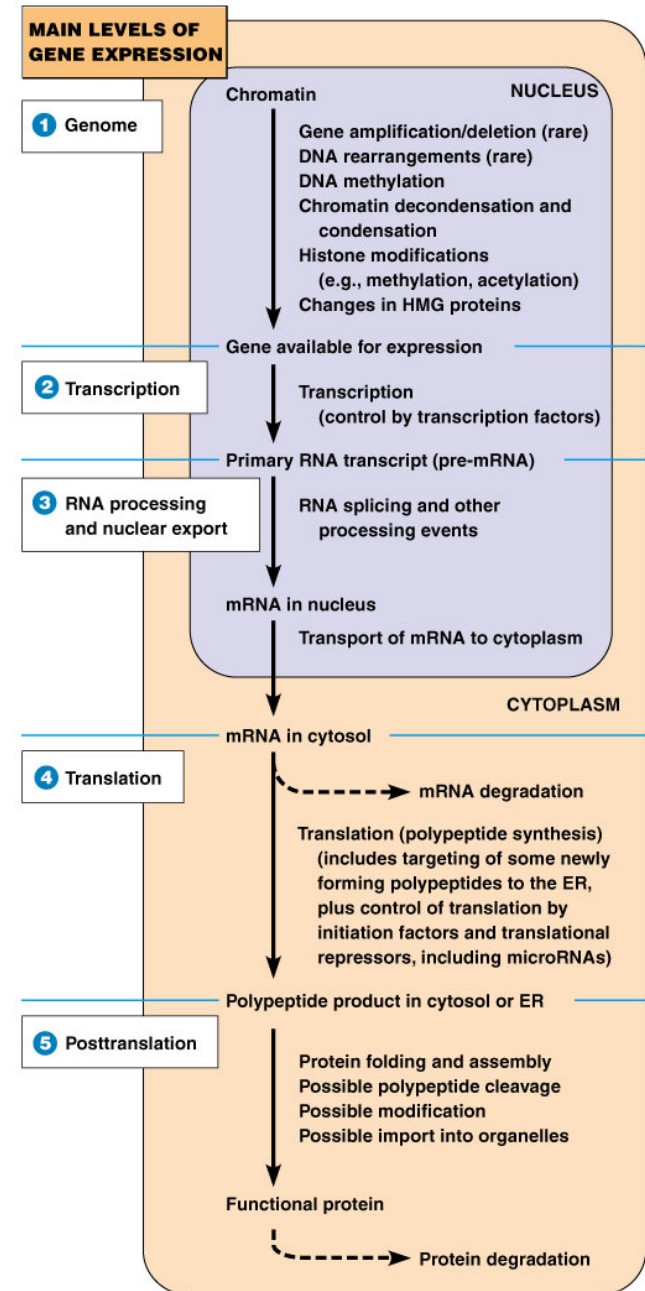
Genome Level (Chromatin Remodelling and DNA rearrangements)

Transcriptional Control

Post-Transcriptional Control

Translational Control

Post-Translational Control (protein activity control)

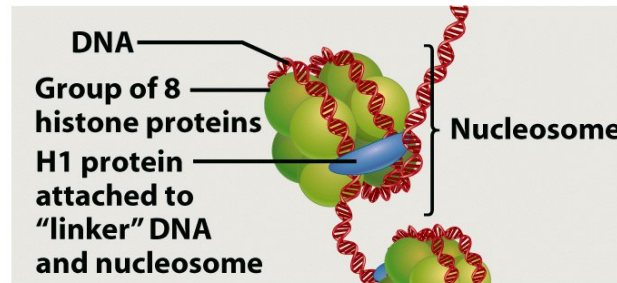




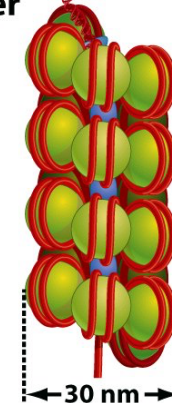
# IV. Gene Regulation in Eukaryotes

## IV.1 Chromatin structure

### (b) Nucleosome structure



### (c) 30-Nanometer fiber



# IV. Gene Regulation in Eukaryotes

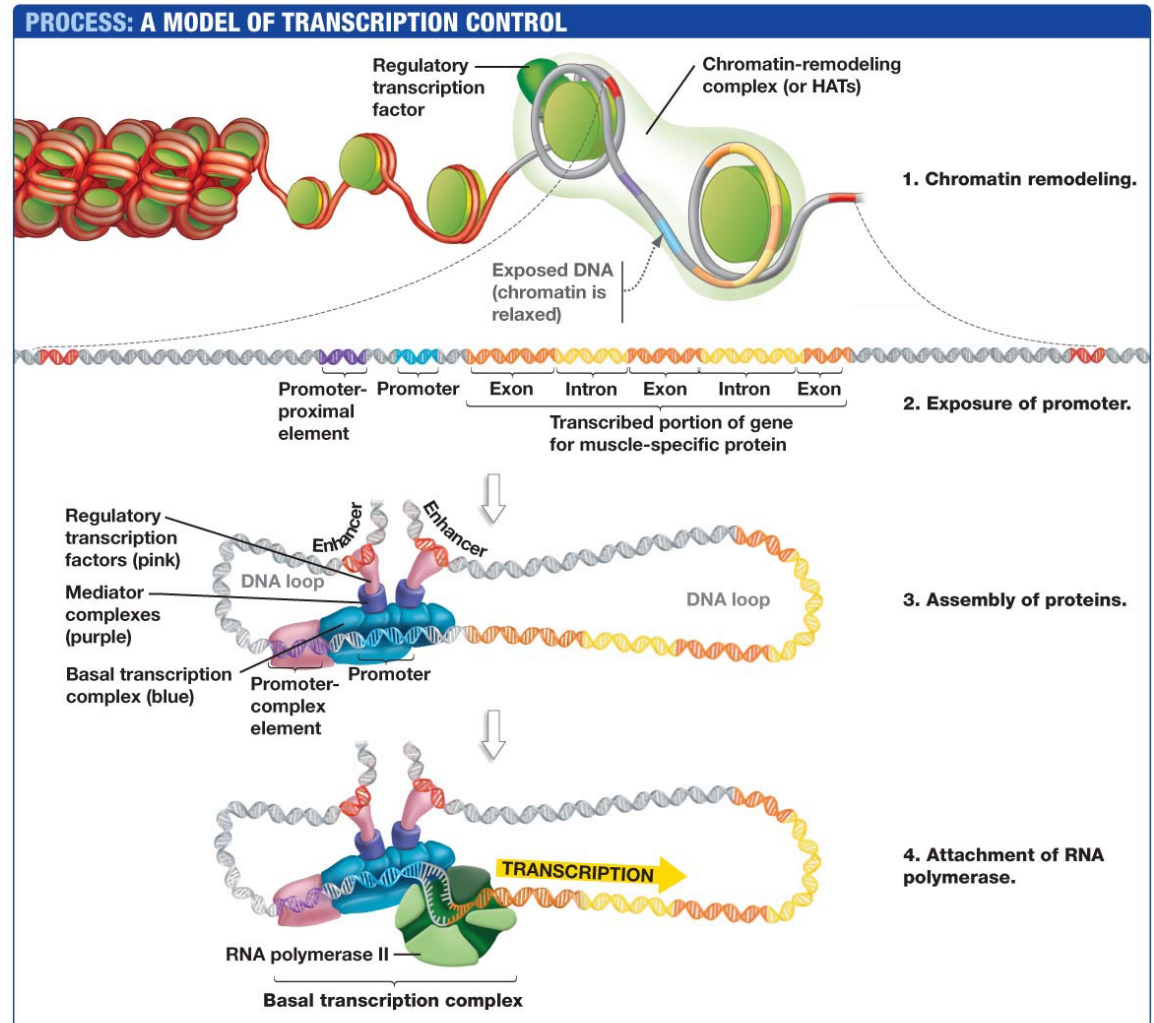
## IV.2 Transcriptional control

Three Classes of Proteins Are Involved in Transcriptional Activation

basal transcription factors (TBP)

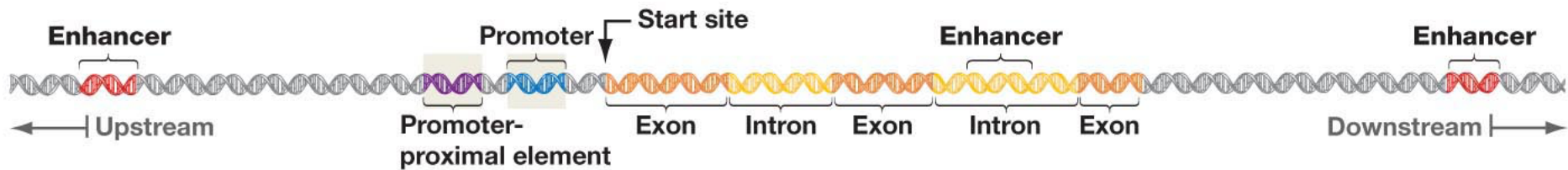
DNA-binding specific TF,

Coactivators (Mediator).



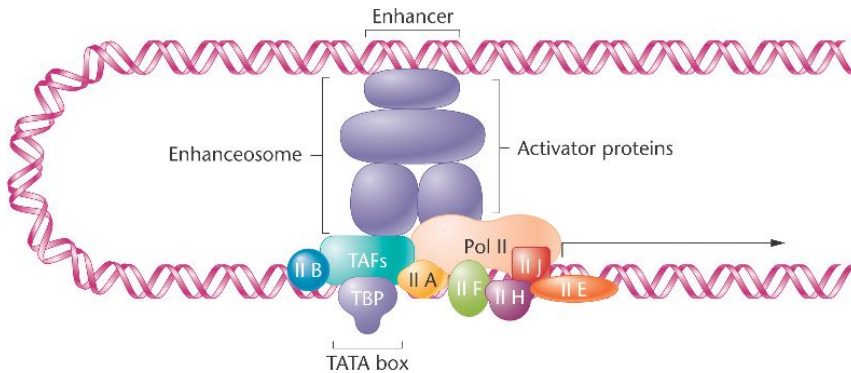
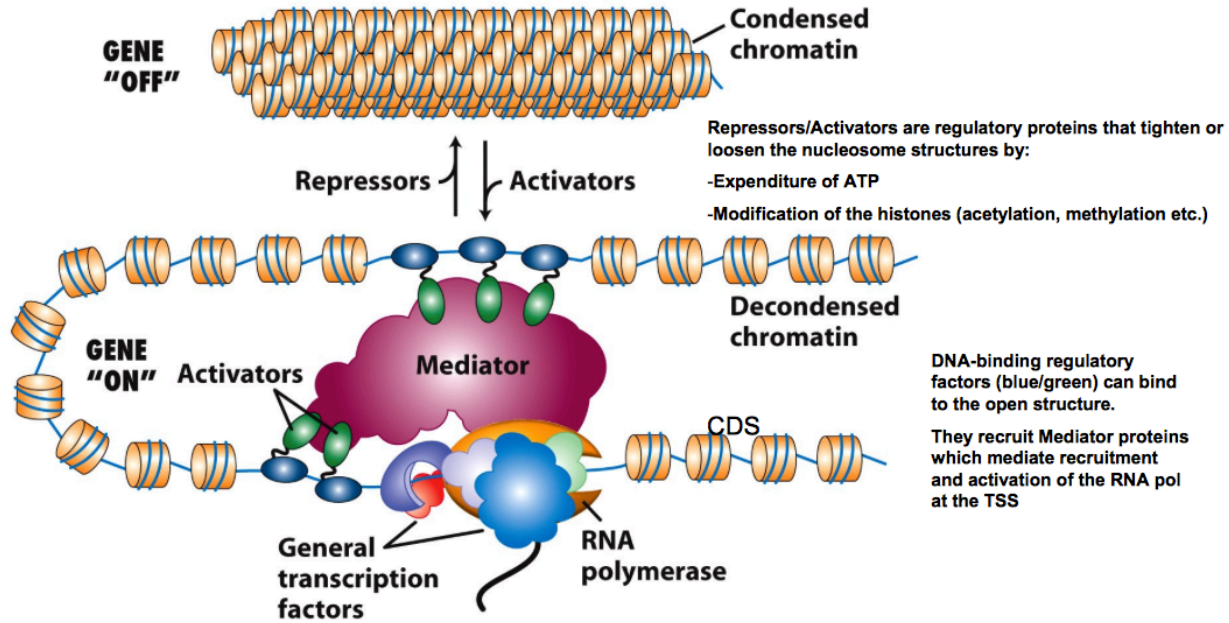
## IV. Gene Regulation in Eukaryotes

Eukaryotic regulatory sequences can be close or far from the core promoter



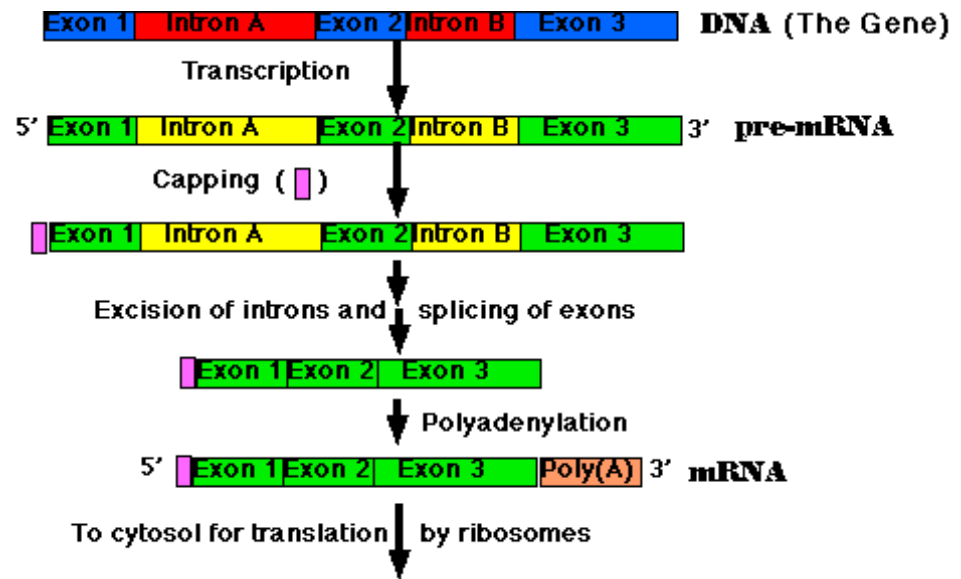
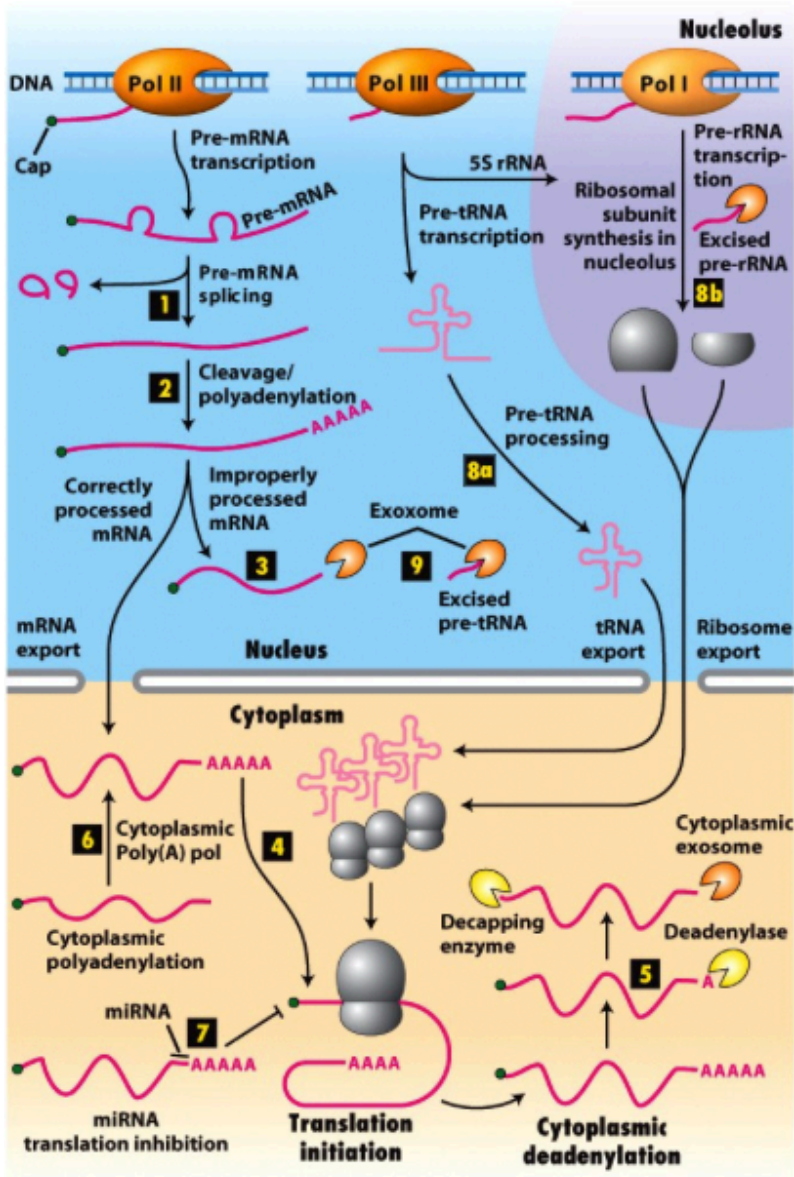
# IV. Gene Regulation in Eukaryotes

Transcription factors are key regulators of eukaryotic gene expression



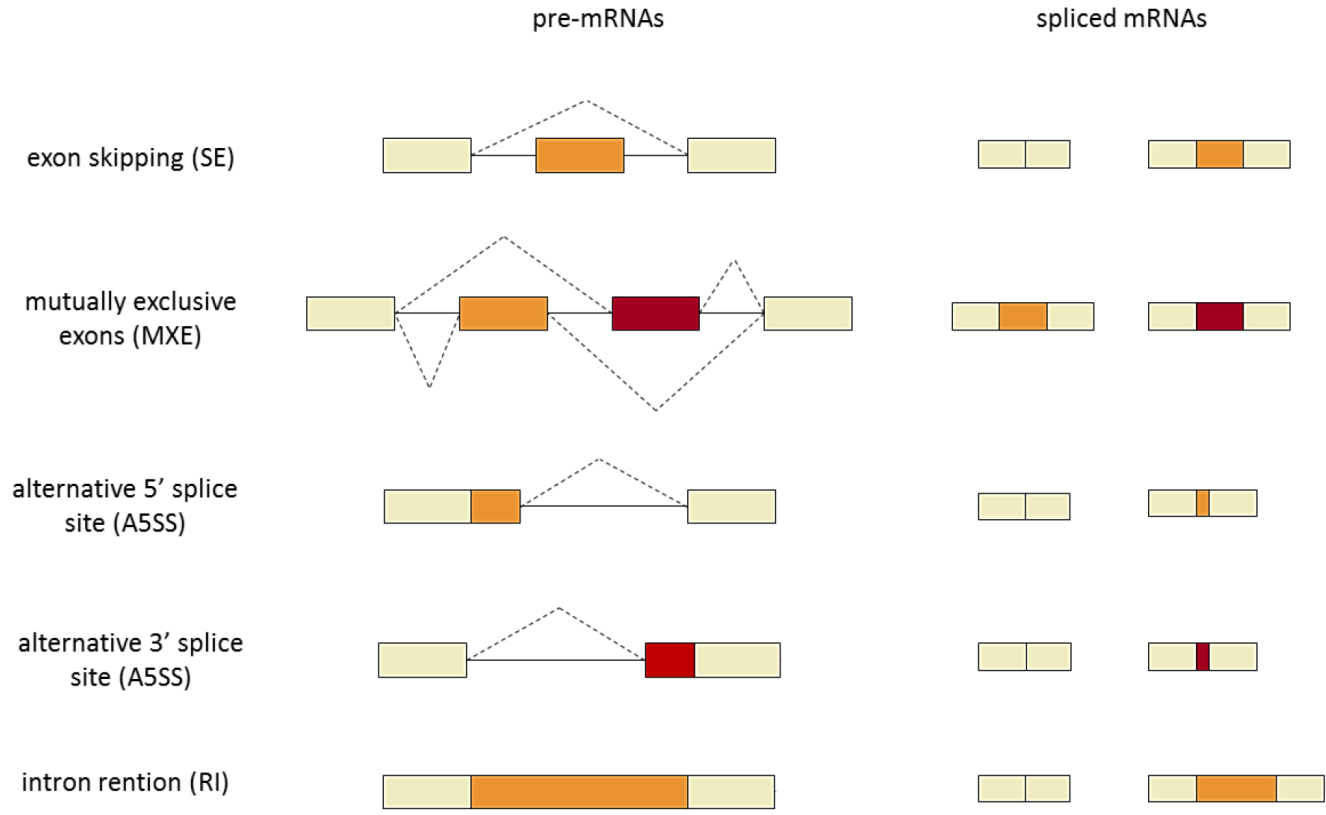
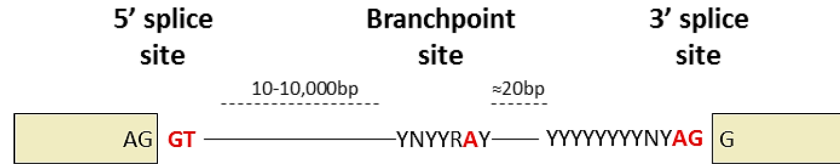
### III. Gene Regulation in Eukaryotes

#### IV.3 Post-transcriptional control of gene expression



### III. Gene Regulation in Eukaryotes

#### IV.3 Post-transcriptional control of gene expression : splicing



## IV. Gene Regulation in Eukaryotes

### IV.3 Post-transcriptional control of gene expression : non coding RNA

only 1% of the genome is translated into proteins but much more transcribed

A great portion of the (**procaryotic and eukaryotic**) genome is transcribed into non coding RNAs. Some of which can be involved in gene regulation :

- sncRNA
- LncRNA
- miRNAs



# IV. Gene Regulation in Eukaryotes

SUMMARY TABLE 18.1 **Regulating Gene Expression in Bacteria and Eukaryotes**

Level of Regulation	Bacteria	Eukaryotes
<b>Chromatin remodeling</b>	<ul style="list-style-type: none"> <li>Limited packaging of DNA</li> <li>Remodeling not a major issue in regulating gene expression.</li> </ul>	<ul style="list-style-type: none"> <li>Extensive packaging of DNA</li> <li>Chromatin must be opened for transcription to begin.</li> </ul>
<b>Transcription</b>	<ul style="list-style-type: none"> <li>Positive and negative control by regulatory proteins that act at sites close to the promoter</li> <li>Sigma interacts with promoter.</li> </ul>	<ul style="list-style-type: none"> <li>Positive and negative control by regulatory proteins that act at sites close to and far from promoter</li> <li>Large basal transcription complex interacts with promoter.</li> <li>Mediator complex required.</li> </ul>
<b>RNA processing</b>	<ul style="list-style-type: none"> <li>None documented</li> </ul>	<ul style="list-style-type: none"> <li>Extensive processing: alternative splicing of introns</li> <li>addition of 5' cap and 3' tail</li> </ul>
<b>mRNA stability</b>	<ul style="list-style-type: none"> <li>Some RNA interference documented</li> </ul>	<ul style="list-style-type: none"> <li>For many genes, RNA interference limits life span or translation rate.</li> </ul>
<b>Translation</b>	<ul style="list-style-type: none"> <li>Regulatory proteins bind to mRNAs and/or ribosome and affect translation rate.</li> </ul>	<ul style="list-style-type: none"> <li>Regulatory proteins bind to mRNAs and/or ribosome and affect translation rate.</li> </ul>
<b>Post-translational modification</b>	<ul style="list-style-type: none"> <li>Folding by chaperone proteins</li> <li>Chemical modification (e.g., phosphorylation) may change activity.</li> </ul>	<ul style="list-style-type: none"> <li>Folding by chaperone proteins</li> <li>Chemical modification (glycosylation, phosphorylation)</li> <li>Ubiquitination targets proteins for destruction by proteasome.</li> </ul>



# V. Principles of transcriptome analysis

## V.1 SAGE Technology

SAGE : Serial analysis of gene expression (1995) :

Unique tags are used to identify each mRNA

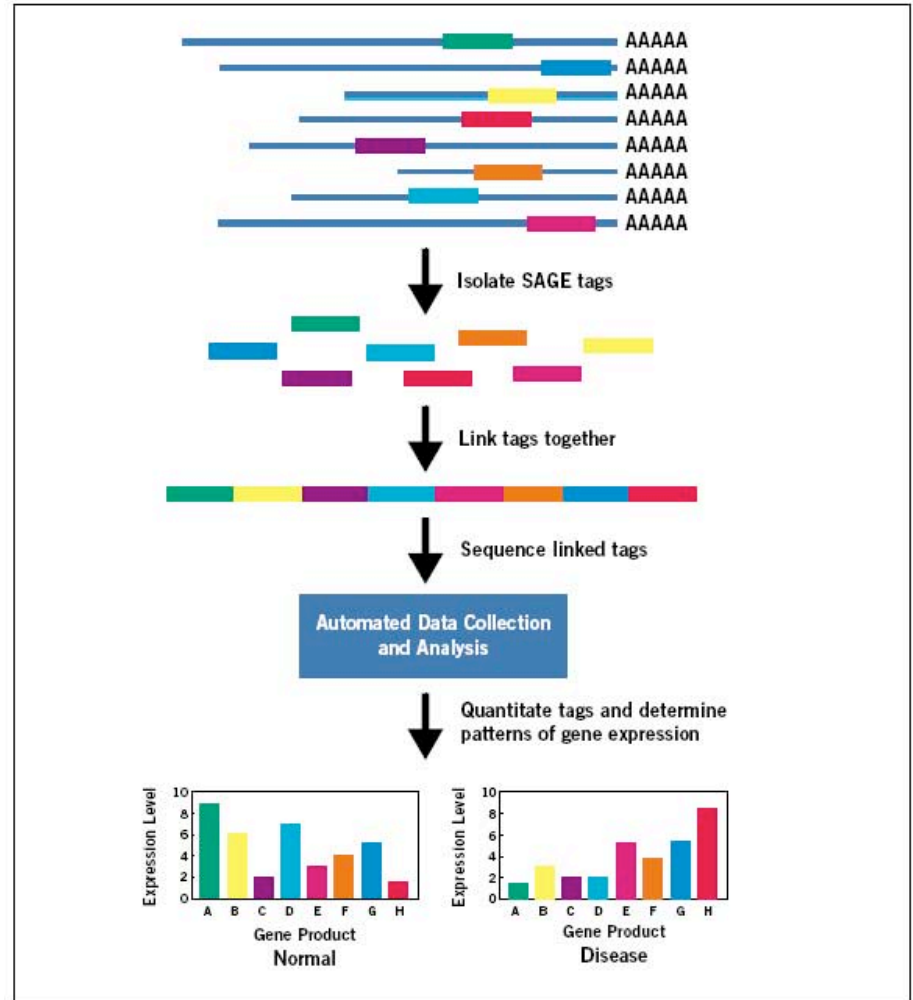
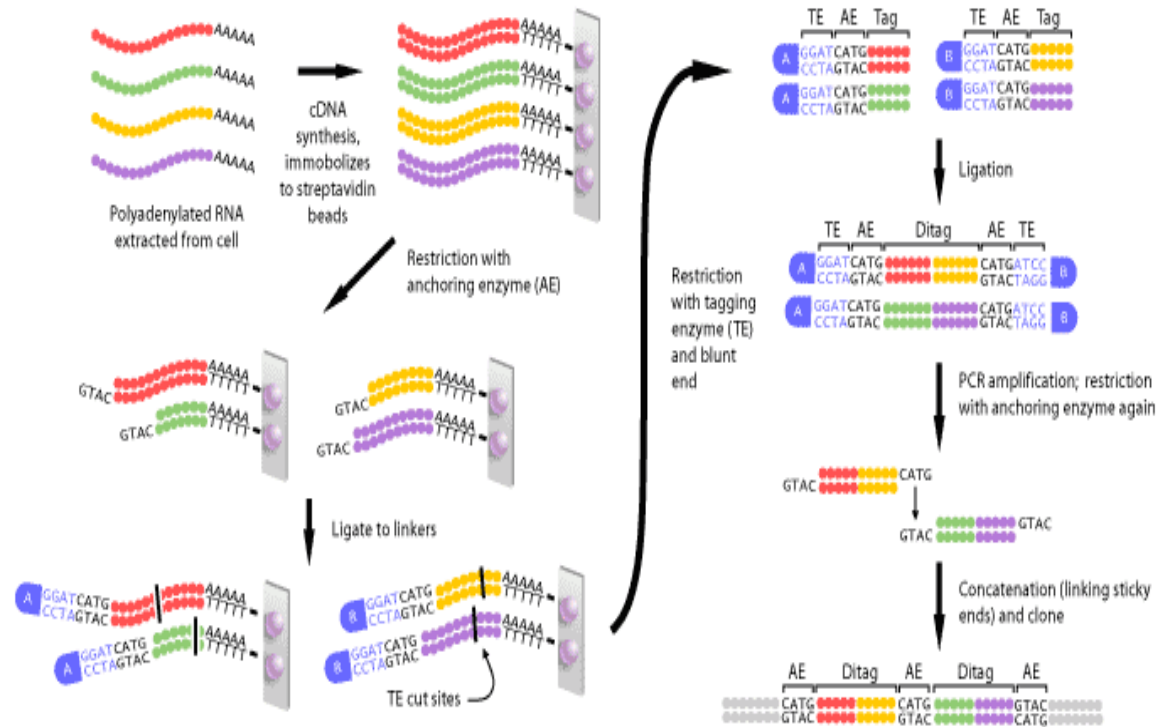


Figure 1. Schematic of SAGE method (Courtesy of sagenet.org)<sup>3</sup>

# V. Principles of transcriptome analysis

## V.1 SAGE Technology

1. Isolate the **mRNA** of an input sample (e.g. a **tumour**). Use Reverse Transcriptase to synthesize complementary DNA (**cDNA**) from mRNA
2. Extract a small chunk of sequence from a defined position of each **cDNA** molecule.
3. Link these small pieces of sequence together to form a long chain (or concatamer).
4. Clone these chains into a **vector** which can be taken up by bacteria.
5. Sequence these chains using modern high-throughput **DNA sequencers**.
6. Process this data with a computer to count the small sequence tags.



# V. Principles of transcriptome analysis

## V.1 SAGE Technology

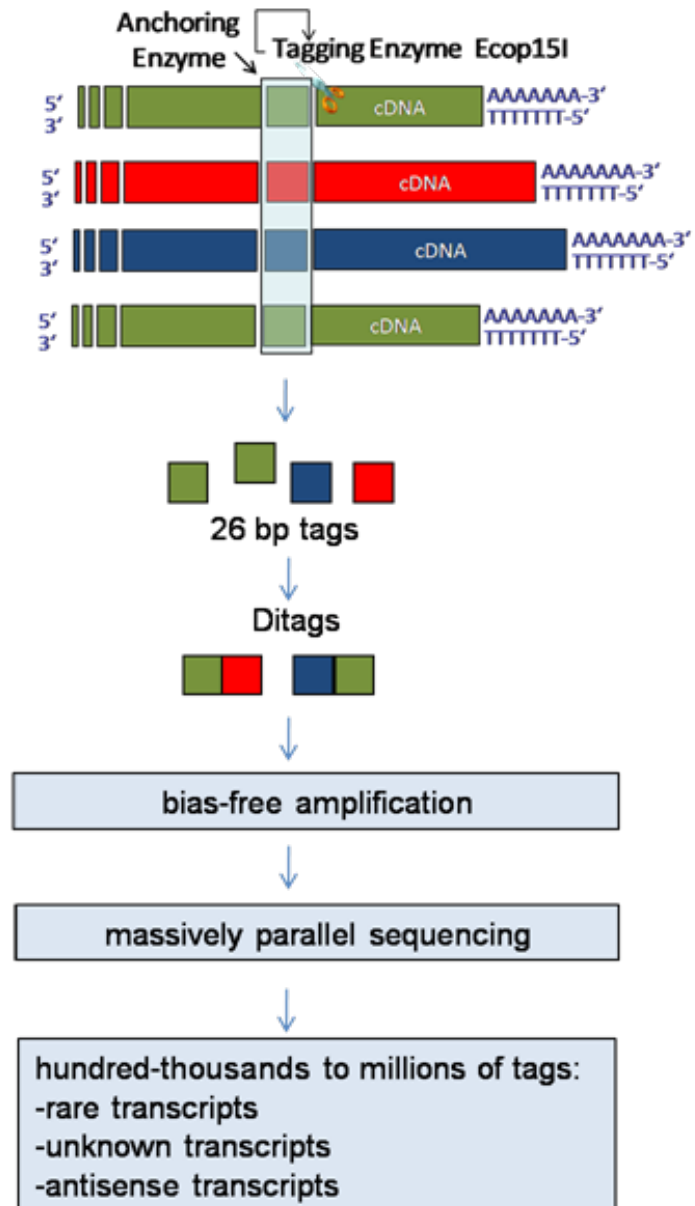
TAG	COUNT	TAG	COUNT	TAG	COUNT
CCCATCGTCC	1286	CACTACTCAC	245	TTCACTGTGA	150
CCTCCAGCTA	715	ACTAACACCC	229	ACGCAGGGAG	142
CTAAGACTTC	559	AGCCCTACAA	222	TGCTCCTACC	140
GCCCAGGTCA	519	ACTTTTTCAA	217	CAAACCATCC	140
CACCTAATTG	469	GCCGGGTGGG	207	CCCCCTGGAT	136
CCTGTAATCC	448	GACATCAAGT	198	ATTGGAGTGC	136
TTCATACACC	400	ATCGTGGCGG	193	GCAGGGCCTC	128
ACATTGGGTG	377	GACCCAAGAT	190	CCGCTGCACT	127
GTGAAACCCC	359	GTGAAACCCCT	188	GGAAAACAGA	119
CCACTGCACT	359	CTGGCCCTCG	186	TCACCGGTCA	118
TGATTTCACT	358	GCTTTATTTG	185	GTGCACTGAG	118
ACCCTTGGCC	344	CTAGCCTCAC	172	CCTCAGGATA	114
ATTTGAGAAG	320	GCGAAACCCCT	167	CTCATAAGGA	113
GTGACCACGG	294	AAAACATTCT	161	ATCATGGGGA	110

- Collect sequence records from GenBank
- Assign sequence orientation (by finding poly-A tail or poly-A signal or from annotations)
- Assign UniGene identifier to each sequence with a SAGE tag
- Record (for each tag-gene pair)
  - #sequences with this tag
  - #sequences in gene cluster with this tag
- Softwares : [SAGE Genie](#) or SAGE map

# V. Principles of transcriptome analysis

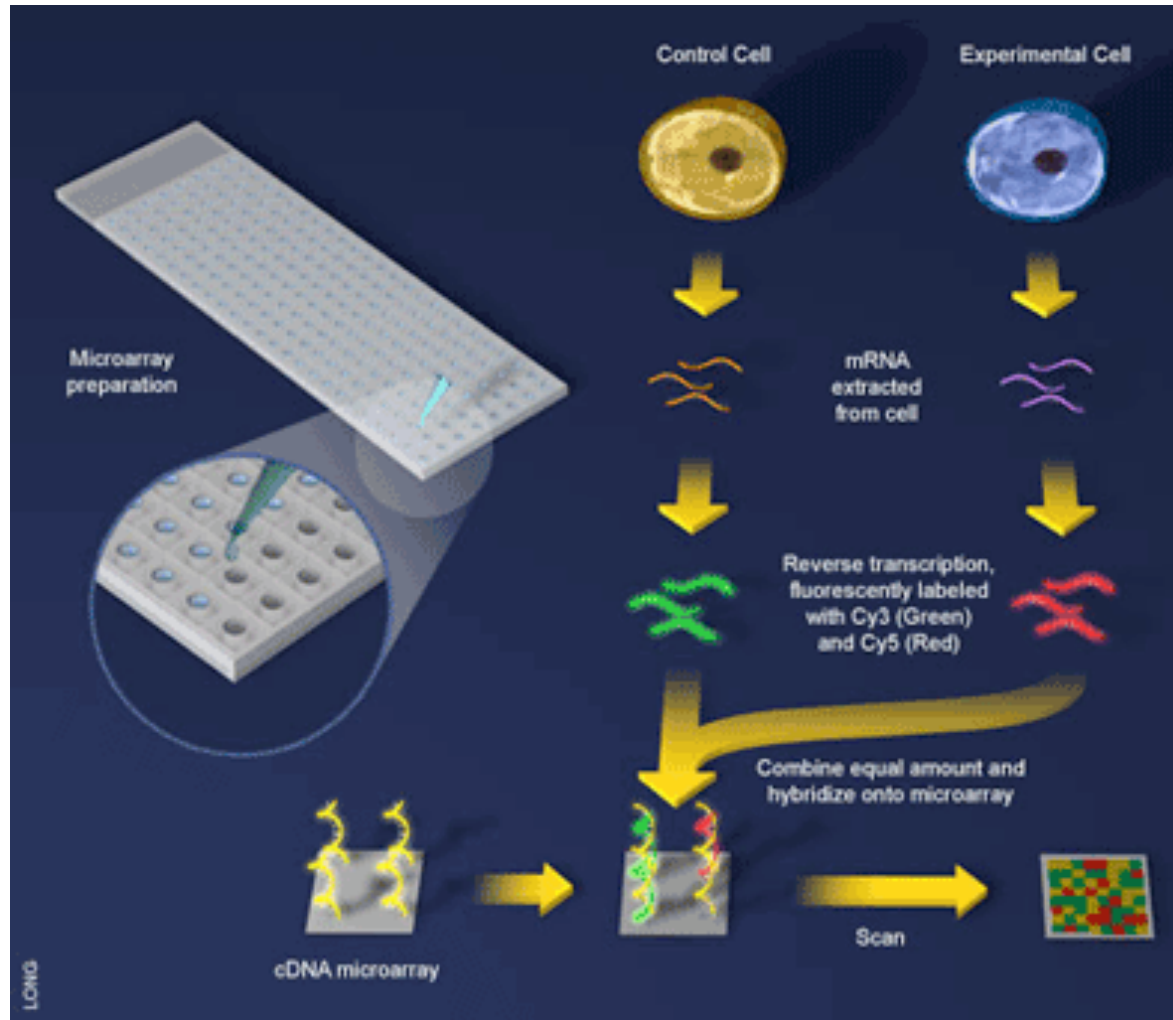
## V.1 SAGE Technology

HT-SuperSAGE



# V. Principles of transcriptome analysis

## V.2 Microarray Technology



[Microarray technology animation](#)

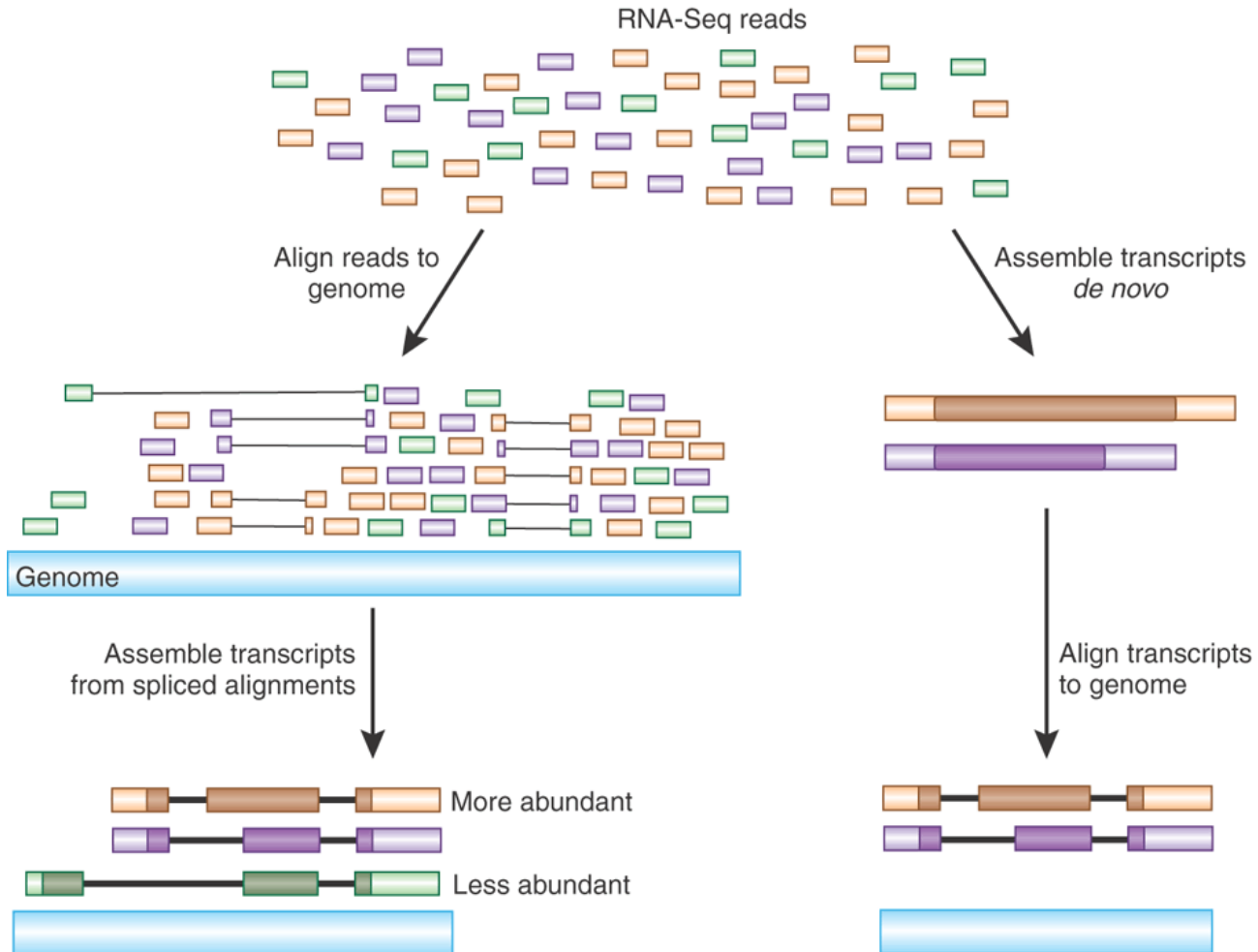
## V. Principles of transcriptome analysis

### V.2 Microarray Technology

<i>Features</i>	<i>SAGE</i>	<i>Microarray</i>
<i>Detects unknown transcripts</i>	Yes	No
<i>Quantification</i>	Absolute measure	Relative measure
<i>Sensitivity</i>	High	Moderate
<i>Specificity</i>	Moderate	High
<i>Reproducibility</i>	Good for higher abundance transcripts	Good for data from intra-platform comparison
<i>Direct cost</i>	5-10X higher than arrays.	5-10 X lower than SAGE

# V. Principles of transcriptome analysis

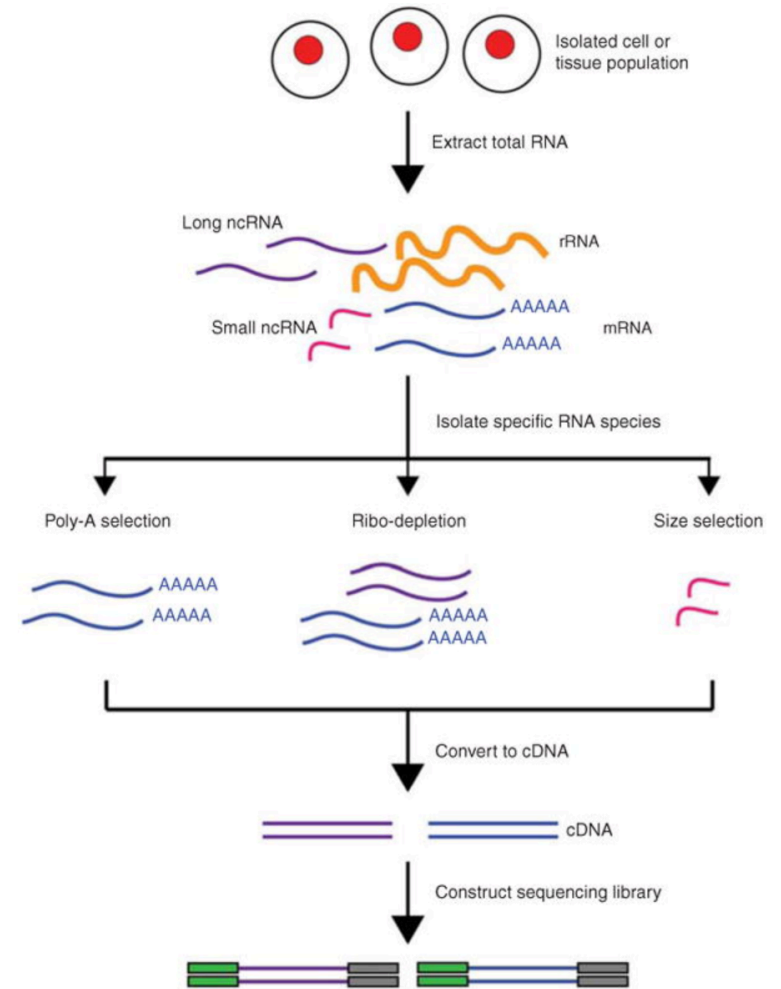
## V.3 RNA-Sequencing (RNA-Seq) : basics



# V. Principles of transcriptome analysis

## V.3 RNA-Sequencing (RNA-Seq) : methodology

1. RNA extraction
2. Reverse transcription
3. Library preparation
4. Polymerase Chain reaction (on solid-Illumina or emulsion-454)
5. Sequencing (reversible terminator or pyrosequencing)
6. Bioinformatic processing

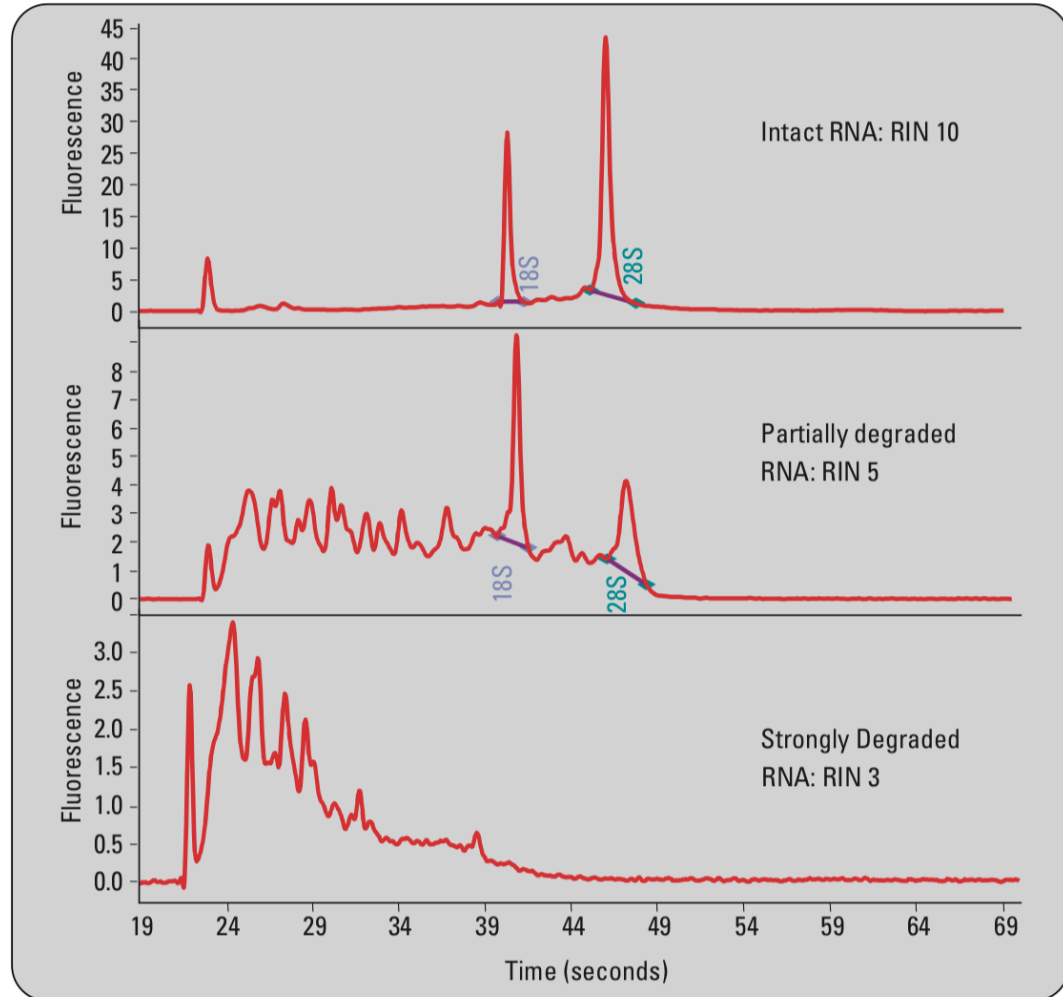
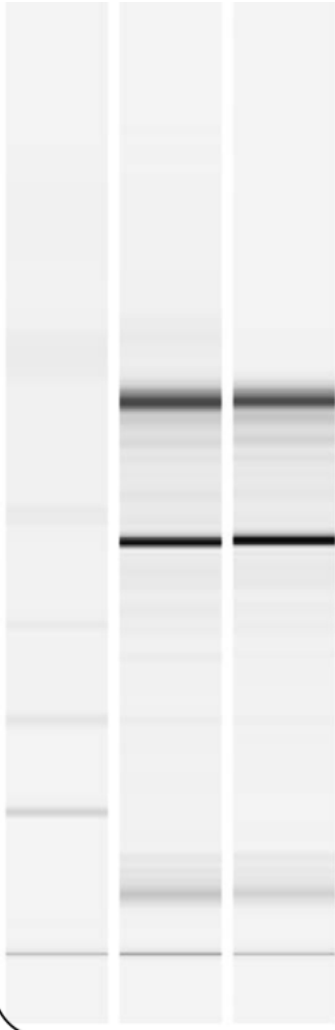




# V. Principles of transcriptome analysis

## V.3 RNA-Sequencing (RNA-Seq) : methodology

### 1. RNA extraction and QC (Agilent Bioanalyzer)



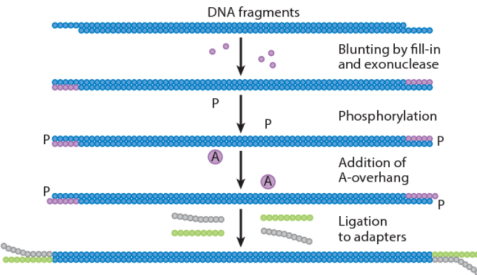
# V. Principles of transcriptome analysis

## V.3 RNA-Sequencing (RNA-Seq) : methodology

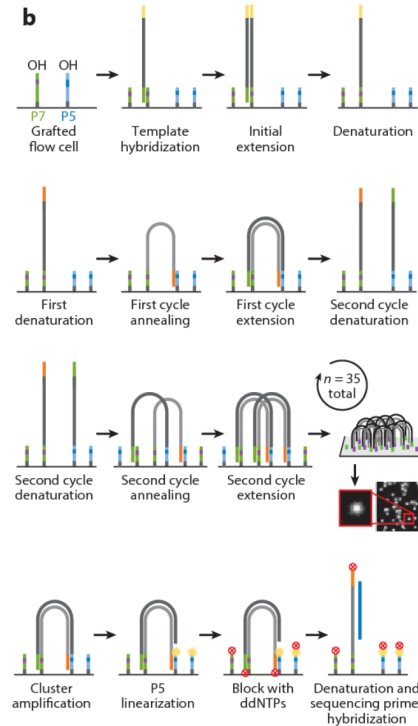
### 2. Reverse transcription, library preparation, amplification, sequencing

#### ILLUMINA

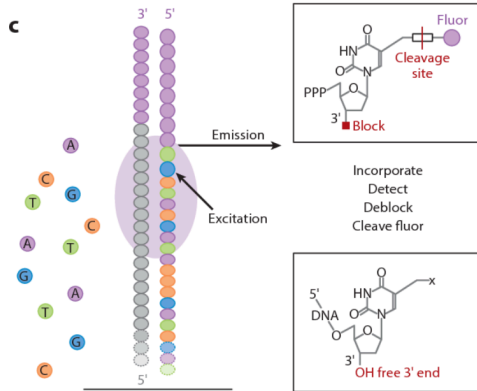
**a** Illumina's library-preparation workflow



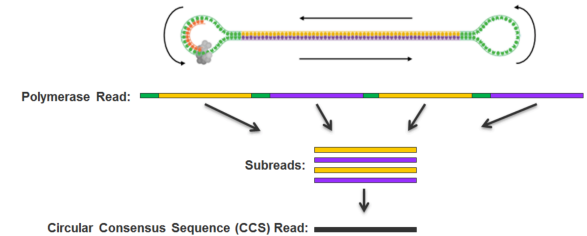
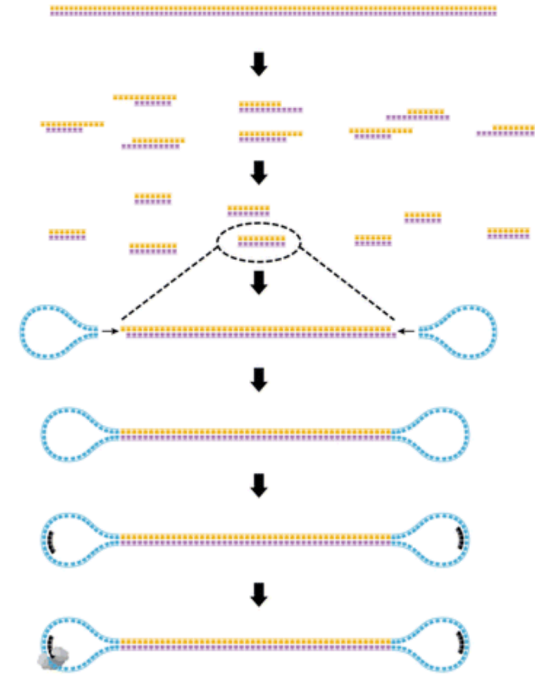
**b**



**c**



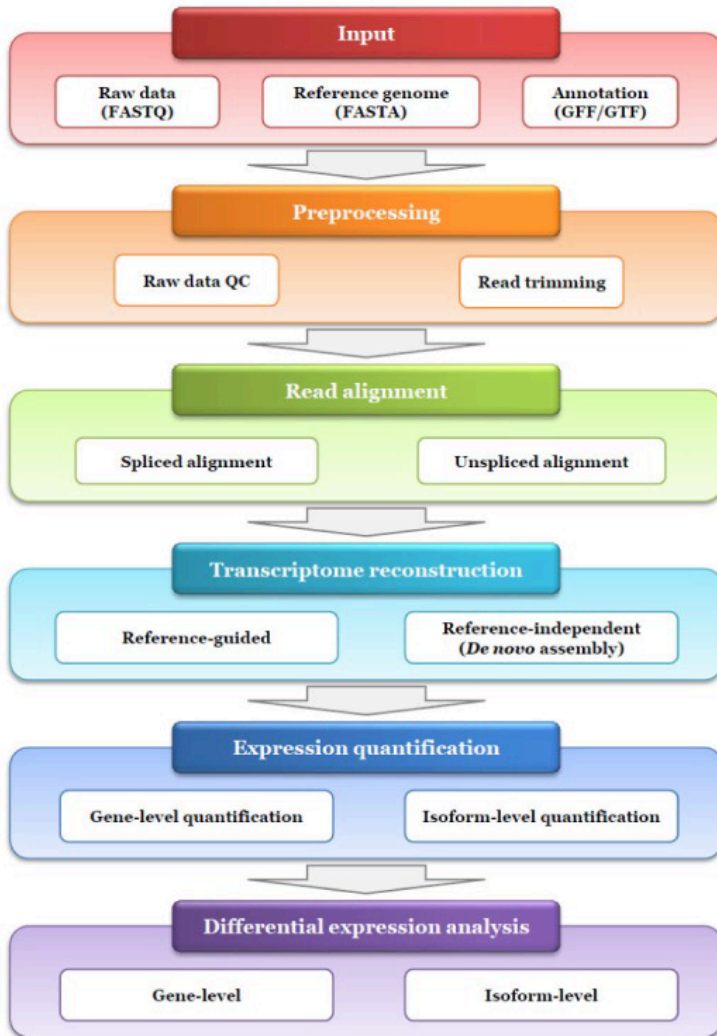
#### PACIFIC BIOSCIENCES



# V. Principles of transcriptome analysis

## V.3 RNA-Sequencing (RNA-Seq) : methodology

### 6. Bio-informatics



Workflow	Category	Package
Preprocessing of raw data	Raw data QC	FastQC
		HTQC
	Read trimming	FASTX-Toolkit
Read alignment	Unspliced aligner	FLEXBAR
		MAQ
		BWA
	Spliced aligner	Bowtie
		TopHat
		MapSplice
RNA-seq specific quality control		STAR
		GSNAP
		RNA-SeQC
		RSeQC
Transcriptome reconstruction	Reference-guided	Qualimap 2
		Cufflinks
		Scripture
	Reference-independent	StringTie
		Trinity
Expression quantification	Gene-level quantification	Oases
		transABySS
		ALEXA-seq
		Enhanced read analysis of gene expression (ERANGE)
		Normalization by expected uniquely mappable area (NEUMA)
		Differential expression
StringTie		
RSEM		
Sailfish		
Differential expression	Gene-level	NOIseq
		edgeR
		DESeq
	Isoform-level	SAMseq
		Cuffdiff
		EBSeq
		Ballgown

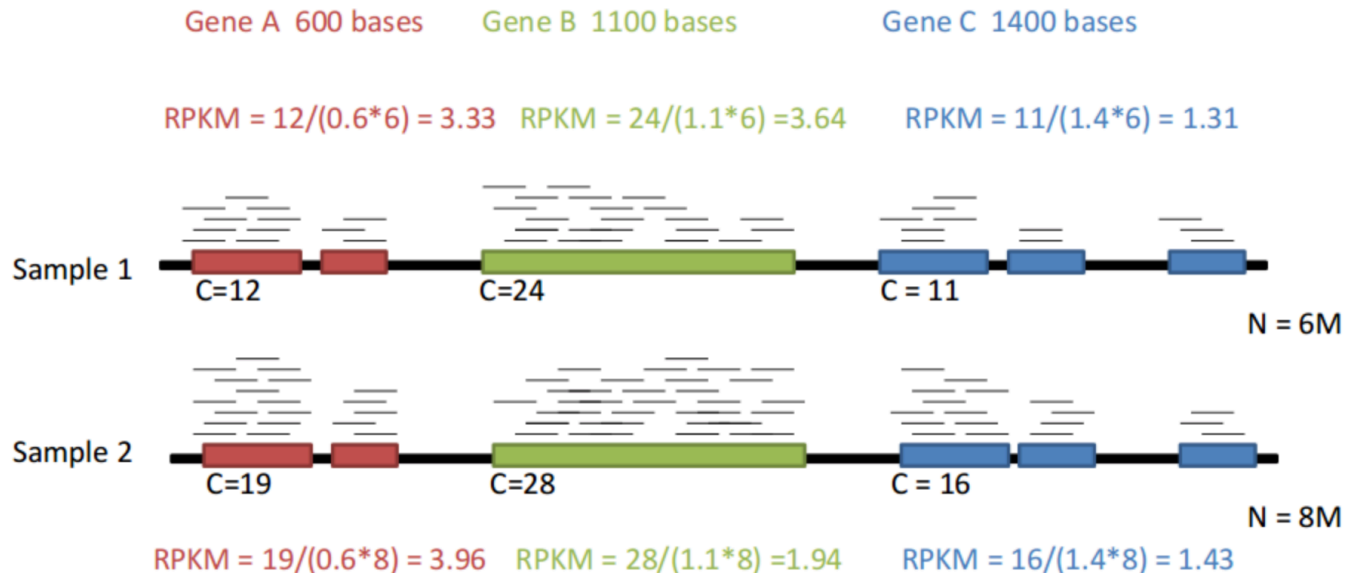
## IV. Principles of transcriptome analysis

### IV.3 RNA-Sequencing (RNA-Seq)

#### Gene expression levels

$$RPKM = \frac{\text{number of reads of the region}}{\frac{\text{total reads}}{1,000,000}} \times \frac{\text{region length}}{1,000}$$

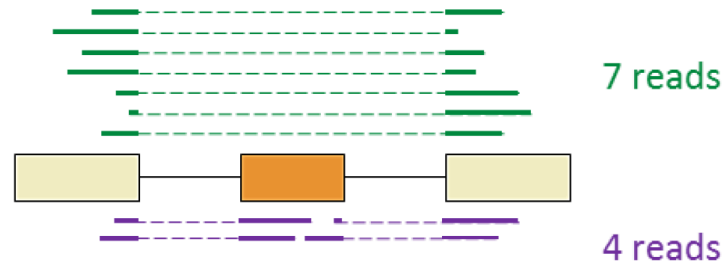
### RPKM Example



## IV. Principles of transcriptome analysis

### IV.3 RNA-Sequencing (RNA-Seq)

#### Splicing analysis



$$\text{PSI} = \frac{\# \text{ inclusion reads}}{(\# \text{ inclusion reads} + \# \text{ skipping reads})} = \frac{4}{(4 + 7)} = 0.36$$

## **IV. Principles of transcriptome analysis**

### **IV.3 RNA-Sequencing (RNA-Seq)**

**Detection of novel transcripts and other changes**

**Dynamic range**

**Specificity and sensitivity**

**Costly and requires high computational power**