

KARPAGAM ACADEMY OF HIGHER EDUCATION

DEPARTMENT OF BIOTECHNOLOGY II B.Sc., BIOTECHNOLOGY SEMESTER III

18BTU302	MOLECULAR BIOLOGY	4H - 4C
Total hours/week: L:4 T:0 P:	Marks: Internal: 40 External: 60	Total: 100

Course Objectives: This course deals with the molecular characterization, structures and functions of cells.

Course Outcomes: The students will enable to know about the theoretical knowledge about the molecular biology and molecular techniques to identify, isolate nucleic acid.

UNIT-I

DNA structure and organization: DNA as genetic material, Structure of DNA, Types of DNA, Organization of DNA in prokaryote and eukaryotic cells, Chromosome biology – histone and non-histone proteins, organization.

UNIT-II

DNA replication: Replication of DNA in prokaryotes and eukaryotes: Semi-conservative nature of DNA replication, Bi-directional replication, DNA polymerases, Replication complex: Pre-priming proteins, primosome, replisome, Rolling circle replication, Unique aspects of eukaryotic chromosome replication, Fidelity of replication.

UNIT-III

Transcription and RNA processing: RNA structure and types of RNA, Transcription in prokaryotes: Prokaryotic RNA polymerase, role of sigma factor, promoter, Initiation, elongation and termination of RNA chains Transcription in eukaryotes: Eukaryotic RNA polymerases, transcription factors, promoters, enhancers, mechanism of transcription initiation, promoter clearance and elongation RNA splicing and processing: processing of pre-mRNA: 5' cap formation, polyadenylation, splicing, rRNA and tRNA splicing.

UNIT-IV

Regulation of gene expression and translation: Regulation of gene expression in prokaryotes: Operon concept (inducible and repressible system), Genetic code and its characteristics, Prokaryotic and eukaryotic translation: ribosome structure and assembly, Charging of tRNA, aminoacyl tRNA synthetases, Mechanism of initiation, elongation and termination of polypeptides, Fidelity of translation, Inhibitors of translation, Posttranslational modifications of proteins.

UNIT-V

DNA damage, repair and homologous recombination: DNA damage and repair: causes and types of DNA damage, mechanism of DNA repair: Photoreactivation, base excision repair, nucleotide excision repair, mismatch repair, trans-lesion synthesis, recombinational repair, nonhomologous end joining. Homologous recombination: models and mechanism.

References

- 1. Karp, G. (2013). *Cell and Molecular Biology: Concepts and Experiments* (7th ed.). Hoboken, US: John Wiley & Sons. Inc.
- 2. Watson, J. D., Baker T.A., Bell, S. P., Gann, A., Levine, M., & Losick, R. (2008). *Molecular Biology of the Gene* (6th ed.). Cold Spring Harbour Lab. Press, Pearson Pub.
- 3. De Robertis, E.D.P., & De Robertis, E.M.F. (2006). *Cell and Molecular Biology* (8th ed.). Lippincott Williams and Wilkins, Philadelphia.
- 4. Becker, W.M., Kleinsmith, L.J., Hardin. J., & Bertoni, G. P. (2009). *The World of the Cell* (7th ed.). San Francisco: Pearson Benjamin Cummings Publishing.



KARPAGAM ACADEMY OF HIGHER EDUCATION

DEPARTMENT OF BIOTECHNOLOGY

II B.Sc., BIOTECHNOLOGY - SEMESTER 3

LECTURE PLAN - MOLECULAR BIOLOGY

18BTU302

S.No	Lecture Duration (hr)	Topic to be covered	Support materials
I	()	UNIT I (Total No. of hours - 7 hours)	
1	1	DNA as Genetic Material	
2	1	Structure of DNA	T2 97-110
3	1	Types of DNA	
4	1	Organization of DNA in Prokaryotes	W1
5	1	Organization of DNA in Eukaryotes	W1
6	1	Chromosome structure – histone and Non-histone proteins	R1 53-65
7	1	Unit test	-
I		UNIT II (Total No. of hours - 10 hours)	
8	1	DNA replication – Introduction	R1 131-135
9	1	Replication of DNA in Prokaryotes	J1 395-420
10	1	Replication of DNA in Eukaryotes	W2
11	1	Semi-conservative nature of DNA replication	T2 109-113
12	1	Bidirectional replication R1 135-	
13	1	DNA Polymerases	T2 109
14	1	Replication Complexes & Pre-priming proteins	T2 158-162
15	1	Rolling Circle replication	T2 149-151
16	1	Fidelity of replication and Unit Revision	T2 143-144
17	1	Unit test -	
		UNIT III (Total No. of hours - 11 hours)	
18	1	RNA structure and types of RNA	R2 60-64
19	1	Transcription in Prokaryotes T2 361-363	

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20	1	Transcription in Eukaryotes	T2 363-366
21	1	RNA Polymerases and role of σ factor	
22	1	Promotor, Initiation, Elongation and termination	
23	1	Eukaryotic RNA Polymerses	R2 315-363
24	1	Transcription factors	
25	1	Mechanism of transcription Initiation	
26	1	RNA splicing and Processing	T2 379
27	1	Processing of different RNA's	T2 412-415
28	1	Unit test	-
		UNIT IV (Total No. of hours - 11 hours)	
29	1	Regulation of gene expression – Overview	T2 410-415
30	1	Regulation of gene expression in Prokaryotes	D2 267 270
31	1	Operon concept – Inducible and repressible	R2 367-378
32	1	Genetic Code and its Characteristics	T1 461-469
33	1	Prokaryotic and Eukaryotic Translation	T1 483-527
34	1	Ribosome structure assembly	R2 443-450
35	1	Charging tRNA, aminoacyl tRNA synthetse	
36	1	Mechanism of Initiation, Elongation and Termination	
37	1	Fidelity of translation	R2 379-396
38	1	Post-translational Modifications	
39	1	Unit test and Discussions	-
		UNIT V (Total No. of hours - 9 hours)	
40	1	DNA Damage – Overview	T1 591-596
41	1	Causes and types of DNA Damage	R2 280-290
42	1	Mechanism of DNA repair	R2 280-290
43	1	Photoreactivation and their genes	R2 280-290
44	1	Base Excision, nucleotide excision and repair mechanism	R2 280-290
45	1	Mismatch repair and translational synthesis	R2 280-290
46	1	Recombinational repair	R2 280-290
47	1	Homologous end joining, Recombination models and Mechanisms	R2 280-290
48	1	Unit test and discussion	-
		Total No. of hours	48 hours
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Reference Books

- T1. Karp, G. (2013). Cell and Molecular Biology: Concepts and Experiments (7th ed). Hoboken, US: John Wiley & Sons. Inc.
- T2. Watson, J.D., Baker T.A., Bell, S.P., Gann, A., Levine, M., & Losick, R. (2008). Molecular Biology of the gene (6th edition). Cold spring Harbour Lab. Press, Pearson Pub.
- 3. R1. Lodish et al. (2006). Molecular Cell Biology (5th edition).
- 4. R2. David Friefielder (2006). Molecular Biology (2006). Narosa Publishing house.
- 5. W1: www.biologydiscussion.com
- 6. W2: <u>www.sciencedirect.com/topics/neuroscience/eukaryotic-dna-replication</u>
- J1: Alberts B.M (1987). Prokaryotic DNA replication Mechanism. Phil.Trans.R. Soc. London. B., 317: 395-420.

MOLECULAR BIOLOGY 18BTU302

UNIT I

DNA structure and organization: DNA as genetic material, Structure of DNA, Types of DNA, Organization of DNA in prokaryote and eukaryotic cells, Chromosome biology – histone and non-histone proteins, organization.

S.No	Lecture Duration	Topic to be covered	Support materials
	(hr)	UNIT I (Total No. of hours - 7 hours)	
	1		
1	1	DNA as Genetic Material	
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7	1	Unit test	-

References

T2. Watson, J.D., Baker T.A., Bell, S.P., Gann, A., Levine, M., &Losick, R. (2008).Molecular Biology of the gene (6th edition).Cold spring Harbour Lab. Press, Pearson Pub.

R2. David Friefielder (2006). Molecular Biology (2006). Narosa Publishing house.

W1: www.biologydiscussion.com

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COURSE CODE: 18BTU302	UNIT: I	BATCH-2018-2021		

DNA as Genetic Material

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All cells have the capability to give rise to new cells and the encoded information in a living cell is passed from one generation to another. The information encoding material is thegenetic or hereditary material of the cell.

The research at the end of the 19th century had verified Mendelian inheritance and it was also believed that the genetic material is in thechromosome.

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However, scientists still didn't know the true features of the genetic material. In the earlytwentieth century, biologists believed that proteins carried genetic information.

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But the Griffith experiment with *Streptococcus pneumoniae*(1928), Avery, MacLeod and McCarty experiment (1944) on transforming principle and Hershey-Chase experiment (1952)on bacteriophage T2, confirms that DNA is genetic material.

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Genetic material is the material that determines the inherited characteristics of a functional organism. It has the following properties:

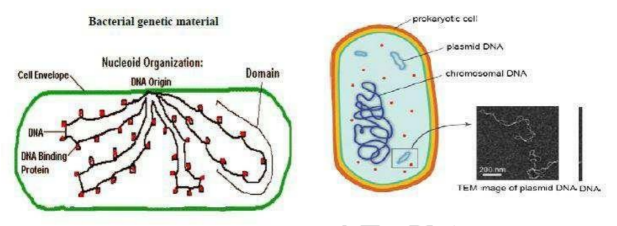
- It must be stable
- It must be capable of being expressed whenneeded
- It must be capable of accuratereplication
- It must be transmitted from parent to progeny withoutchange

Prokaryotic genetic material

- 1. The prokaryotic (bacterial) genetic material is usually concentrated in a specific clear region of the cytoplasm callednucleiod.
- 2. The bacterial chromosome is a single, circular, double stranded DNA molecule mostly attached to the plasma membrane at one point. It does not contain any histone protein. *Escherichia coli* DNA is circular molecule 4.6 million base pairs in length, containing 4288 annotated protein-coding genes (organized into 2584 operons), seven ribosomal RNA (rRNA) operons, and 86 transfer RNA (tRNA) genes. Certain bacteria like the *Borreliaburgdorferi*possess array of linear chromosome likeeukaryotes.
- 3. Besides the chromosomal DNA many bacteria may also carry extra chromosomal genetic elements in the form of small, circular and closed DNA molecules, calledplasmids. They

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generally remain floated in the cytoplasm and bear different genes based on which they have been studied. Some of the different types of plasmids are F plasmids, R plasmids, virulent plasmids, metabolic plasmids etc. Below Figure depicts a bacterial chromosome and plasmid.



Virus genetic material

- 1. The chromosomal material of viruses is DNA or RNA which adopts different structures.
- 2. It is circular when packaged inside the virusparticle.

Eukaryotic genetic material

- 1. A Eukaryotic cell has genetic material in the form of genomic DNA enclosed within the nucleus.
- 2. Genes or the hereditary units are located on the chromosomes which exist as chromatin network in the non-dividing cell/interphase. This will be discussed in detail in the coming sections.

The Griffith Experiment - Hereditary Information Can Pass between Organisms

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The identification of the nucleus as the repository of hereditary information focused attention on the chromosomes, which were already suspected to be the vehicles of Mendelian

inheritance.

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Specifically, biologists wondered how the **genes**, the units of hereditary information studied by Mendel, were actually arranged in thechromosomes.

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They knew that chromosomes contained both protein and deoxyribonucleic acid (DNA). Whichoftheseheldthegenes?Startinginthelate1920sandcontinuingforabout30years,a

series of investigations addressed this question.

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In 1928, British microbiologist Frederick Griffith made a series of unexpected observations while experimenting with pathogenic (disease-causing) bacteria. When he infected mice with

a virulent strain of *Streptococcus pneumonia* bacteria (then known as *Pneumococcus*), the mice died of bloodpoisoning.

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However, when he infected similar mice with a mutant strain of *S. pneumoniae*that lacked the virulent strain's polysaccharide coat, the mice showed no ill effects. The coatwas

apparently necessary for virulence. The normal pathogenic form of this bacterium is referred to as the S form because it forms smooth colonies on a culture dish. The mutant form, which lacks an enzyme needed to manufacture the polysaccharide capsule, is called the R form because it forms roughcolonies.

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To determine whether the polysaccharide coat itself had a toxic effect, Griffith injected dead bacteria of the virulent S strain into mice; the mice remained perfectly healthy.

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As a control, he injected mice with a mixture containing dead S bacteria of the virulent strain and live coatless R bacteria, each of which by itself did not harm the mice. Unexpectedly, the mice developed disease symptoms and many of them died. The blood of the dead mice was

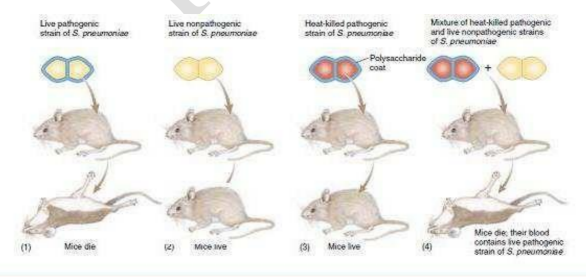
found to contain high levels of live, virulent Streptococcus type S bacteria, which had surface

proteins characteristic of the live (previously R) strain.

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Somehow, the information specifying the polysaccharide coat had passed from the dead, virulent S bacteria to the live, coatless R bacteria in the mixture, permanently transforming

the coatless R bacteria into the virulent S variety. **Transformation** is the transfer of genetic material from one cell to another and can alter the genetic makeup of the recipientcell.



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Griffith's discovery of transformation -

(1) The pathogenic of the bacterium *Streptococcus pneumoniae*kills many of the mice it is injected into. The bacterial cells are covered with a polysaccharide coat, which the bacteria themselvessynthesize.

(2) Interestingly, an injection of live, coatless bacteria produced no ill effects. However, the coat itself is not the agent of disease.

(3) When Griffith injected mice with dead bacteria that possessed polysaccharide coats, the mice were unharmed. (4) But when Griffith injected a mixture of dead bacteria with polysaccharide coats and live bacteria without such coats, many of the mice died, and virulent bacteria with coats were recovered. Griffith concluded that the live cells had been —transformed by the dead ones; that is, genetic information specifying the polysaccharide coat had passed from the dead cells to the livingones.

The Avery Experiments

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The agent responsible for transforming *Streptococcus* went undiscovered until 1944. In a classic series of experiments, Oswald Avery and his coworkers Colin MacLeod and Maclyn

McCarty characterized what they referred to as the -transforming principle.

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They first prepared the mixture of dead S *Streptococcus* and live R *Streptococcus* that Griffith had used. Then Avery and his colleagues removed as much of the protein asthey

could from their preparation, eventually achieving 99.98% purity. Despite the removal of

nearly all protein, the transforming activity was not reduced.

Moreover, the properties of the transforming principle resembled those of DNA in several ways:

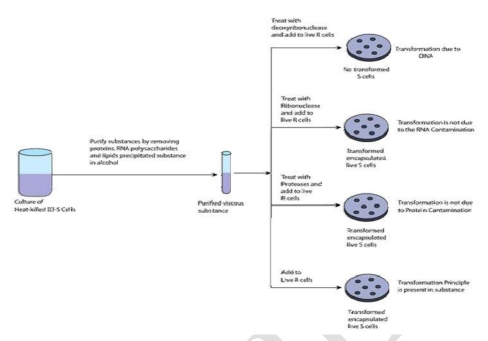
1. When the purified principle was analyzed chemically, the array of elements agreed closely with DNA.

2. When spun at high speeds in an ultracentrifuge, the transforming principle migrated to the same level (density) asDNA.

3. Extracting the lipid and protein from the purified transforming principle did not reduce its activity.

4. Protein-digesting enzymes did not affect the principle's activity; nor did RNA-digesting enzymes.

5. The DNA-digesting enzyme DNase destroyed all transforming activity. The evidence was overwhelming. They concluded that —a nucleic acid of the deoxyribose type is the fundamental unit of the transforming principle of *Pneumococcus* Type IIII—in essence, that DNA is the hereditary material.



Avery, Macleod and McCarty's experiment demonstrating the Griffith's transformation

The Hershey–Chase Experiment

Avery's results were not widely accepted at first, as many biologists preferred to believe that proteins were the repository of hereditary information. Additional evidence supporting

Avery's conclusion was provided in 1952 by Alfred Hershey and Martha Chase, who experimented with **bacteriophages**, viruses that attack bacteria.

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Viruses, consist of either DNA or RNA (ribonucleic acid) surrounded by a protein coat. When a *lytic* (potentially cell-rupturing) bacteriophage infects a bacterialcell, it first binds to

the cell's outer surface and then injects its hereditary information into the cell. There, the hereditary information directs the production of thousands of new viruses within the bacterium. The bacterial cell eventually ruptures, or lyses, releasing the newly madeviruses.

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To identify the hereditary material injected into bacterial cells at the start of an infection, Hershey and Chase used the bacteriophage T2, which contains DNA rather than RNA. They

labeled the two parts of the viruses, the DNA and the protein coat, with different radioactive isotopes that would serve as tracers.

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In some experiments, the viruses were grown on a medium containing an isotope of phosphorus, 32P, and the isotope was incorporated into the phosphate groups of newly

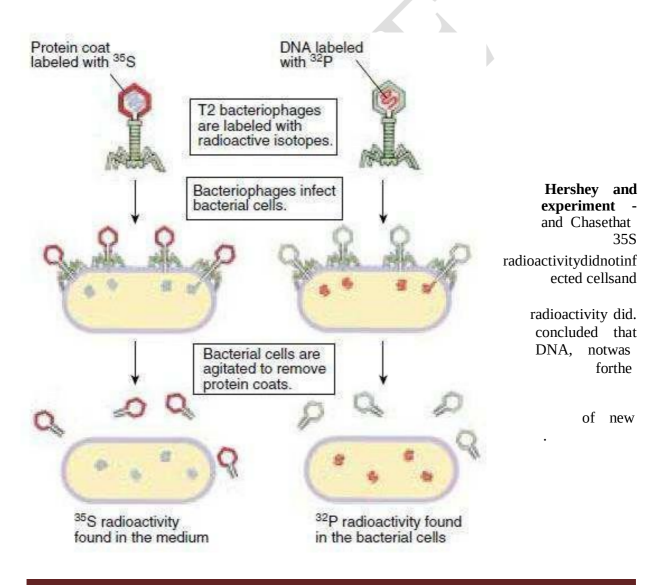
synthesized DNA molecules. In other experiments, the viruses were grown on a medium

containing 35S, an isotope of sulfur, which is incorporated into the amino acids of newly synthesized protein coats. The 32P and 35S isotopes are easily distinguished from eachother because they emit particles with different energies when they decay.

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After the labeled viruses were permitted to infect bacteria, the bacterial cells were agitated violently to remove the protein coats of the infecting viruses from the surfaces of the

bacteria. This procedure removed nearly all of the 35S label (and thus nearly all of the viral protein) from the bacteria. However, the 32P label (and thus the viral DNA) had transferred to the interior of the bacteria and was found in viruses subsequently released from the infected bacteria. Hence, the hereditary information injected into the bacteria that specified the new generation of viruses was DNA and notprotein.



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Different forms of DNA

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The Watson-Crick structure is also referred to as B-form DNA, or B-DNA. The B form is the most stable structure for a random-sequence DNA molecule under physiological

conditions and is therefore the standard point of reference in any study of the properties of DNA. Two structural variants that have been well characterized in crystal structures are the A and Zforms.

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These three DNA conformations are shown, with a summary of their properties. The A form is favored in many solutions that are relatively devoid of water. The DNA is still

arranged in a right-handed double helix, but the helix is wider and the number of base pairs per helical turn is 11, rather than 10.5 as in BDNA.

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The plane of the base pairs in A-DNA is tilted about 20° with respect to the helix axis. These structural changes deepen the major groove while making the minor grooves

hallower. The reagents used to promote crystallization of DNA tend to dehydrate it, and thus most short DNA molecules tend to crystallize in the A form.

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Z-form DNA is a more radical departure from the B structure; the most obvious distinction is the left handed helical rotation. There are 12 base pairs per helical turn, and

the structure appears more slender and elongated. The DNA backbone takes on a zigzag appearance. Certain nucleotide sequences fold into left-handed Z helices much more readily than others. Prominent examples are sequences in which pyrimidines alternate with purines, especially alternating C and G or 5-methyl-C and G residues. To form the left handed helix in Z-DNA, the purine residues flip to the syn conformation, alternating with pyrimidines in the anti-conformation. The major groove is barely apparent in Z-DNA, and the minor groove is narrow anddeep.

Whether A-DNA occurs in cells is uncertain, but there is evidence for some short stretches(tracts)ofZ-DNAinbothbacteriaandeukaryotes'TheseZDNAtractsmayplay

a role (as yet undefined) in regulating the expression of some genes or in genetic recombination.

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	A form	B form	Z form			
Helical sense Diameter	Right handed ~26 Å	Right handed ~20 Å	Left handed ~18 Å			2
Base pairs per helical turn	11	10.5	12			X
Helix rise per base pair	2.6 Å	3.4 Å	3.7 Å		-	2
Base tilt normal to the helix axis	20°	6°	7.			
Sugar pucker conformation	C-3' endo	C-2' endo	C-2' endo for pyrimidines; C-3' endo for purines	 28 Å		
Glycosyl bond conformation	Anti	Anti	Anti for pyrimidines; syn for purines	1		É
					A form	B form

Comparison of A, B, and Z forms of DNA. Each structure shown here has 36 base pairs. The bases are shown in gray, the phosphate atoms in yellow, and the ribose and phosphate oxygens in Blue. Blue is the color used to represent DNA strand is in later chapters. The table summarizes some properties of the three forms of DNA.

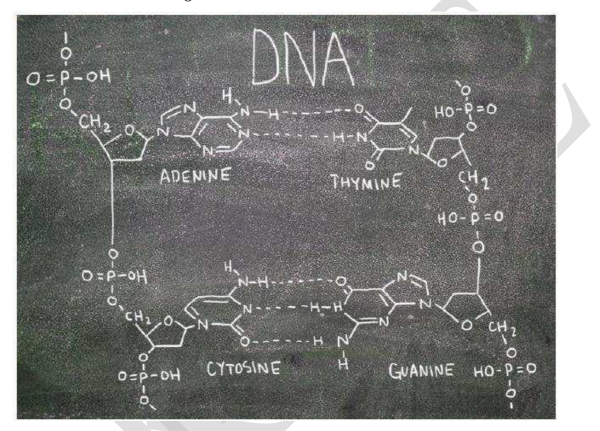
Chargaff's Rules

It was known that DNA is composed of nucleotides, each of which contains a nitrogencontaining base, a five carbon sugar (deoxyribose), and a phosphate group. In these nucleotides, there is one of the four possible bases: adenine (A), guanine (G), cytosine (C), or thymine (T). Adenine and guanine are purine bases, and cytosine and thymine are pyrimidine bases. Erwin Chargaff (1905-2002), an Austrian-American biochemist from Columbia University, analyzed the base composition of the DNA of various species. This led him to propose two main rules that have been appropriately named Chargaff⁴srules.

Rule 1 - Chargaff determined that in DNA, the amount of one base, a purine, always approximately equals the amount of a particular second base, a pyrimidine. Specifically, that in any double-stranded DNA the number of guanine unit sequals approximately the the number of cytosine units and the number of adenine units equals approximately the number of thymine units. Human DNA is 30.9% A and 29.4% T, 19.9% G and 19.8% C. The rule constitutes the basisofbasepairsintheDNAdoublehelix:AalwayspairswithT,andGalwayspairswithC.

He also demonstrated that the number of purines (A+G) always approximates the number of pyrimidines (T+C), an obvious consequence of the base-pairing nature of the DNA double helix.

Rule 2 - In 1947 Chargaff showed that the composition of DNA, in terms of the relative amounts of the A, C, G and T bases, varied from one species to another. This molecular diversity added evidence that DNA could be the genetic material.



Watson and Crick model

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The determination of the structure of DNA by Watson and Crick in 1953 is often said to mark the birth of modern molecular biology. The **Watson–Crick structure** of DNA isof

such importance because, in addition to providing the structure of what is arguably the central molecule of life, it suggested the molecular mechanism of heredity. Watsonand

Crick's accomplishment, which is ranked as one of science's major intellectual achievements, tied together the less than universally accepted results of several diverse studies.

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Information that DNA is a helical molecule. This was provided by an X-ray diffraction photograph of a DNA fiber taken by Rosalind Franklin (DNA, being a threadlike molecule,

does not crystallize but, rather, can be drawn out in fibers consisting of parallel bundles of molecules).

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This photograph enabled Crick, an X-ray crystallographer by training who had earlier derived the equations describing diffraction by helical molecules, to deduce (a) that DNA is a

helical molecule and (b) that its planar aromatic bases form a stack of parallel rings which is parallel to the fiber axis.

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This information only provided a few crude landmarks that guided the elucidation of the DNA structure. It mostly sprang from Watson and Crick's imaginations through model building studies. Once the Watson–Crick model had been published, however, itsbasic simplicity combined with its obvious biological relevance led to its rapid acceptance. Later investigations have confirmed the essential correctness of the Watson–Crick model, although its details have beenmodified.

The Watson–Crick Structure: B-DNA

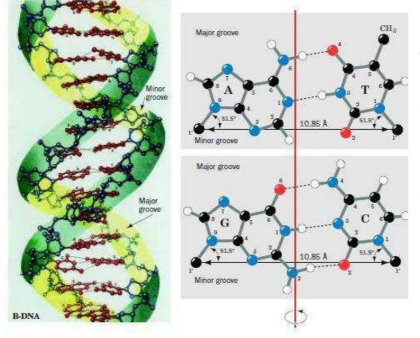
Fibers of DNA assume the so-called B conformation, as indicated by their X-ray diffraction patterns, when the counterion is an alkali metal such as Na1 and the relative humidity is .92%. **B-DNA** is regarded as the **native** (biologically functional) form of DNA because, for example, its X-ray pattern resembles that of the DNA in intact sperm heads. The Watson–Crick structure of B-DNA has the following major features:

1. It consists of two polynucleotide strands that wind about a common axis with a right-handed twist to form an ,20-Å-diameter double helix. The two strands are antiparallel (run in opposite directions) and wrap around each other such that they cannot be separated without unwinding the helix. The bases occupy the core of the helix and the sugar–phosphate chains are coiled about its periphery, thereby minimizing the repulsions between charged phosphategroups.

2. The planes of the bases are nearly perpendicular to the helix axis. Each base is hydrogen bonded to a base on the opposite strand to form a planar base pair. It is these hydrogen bonding interactions, a phenomenon known as **complementary base pairing**, that result in the specific association of the two chains of the doublehelix.

3. The —ideal B-DNA helix has 10 base pairs (**bp**) per turn (a helical twist of 36° per bp) and, since the aromatic bases have van der Waals thicknesses of 3.4 Å and are partially stackedon MOLECULAR BIOLOGY (18BTU302) – Mr. D. Jeyapragash, Asst Prof, Department of Biotechnology, KAHE Page 11

each other (**base stacking**), the helix has a **pitch** (rise per turn) of 34 Å. The most remarkable feature of the Watson–Crick structure is that it can accommodate only two types of base pairs: Each adenine residue must pair with a thymine residue and vice versa, and each guanine residue must pair with a cytosine residue and vice versa. The geometries of these A? T and G?C base pairs, the so-called Watson-Crick base pairs. It can be seen that both of these base pairs are interchangeable in that they can replace each other in the double helix without altering the positions of the sugar-phosphate backbone's C1_i, atoms. Likewise, the double helix is undisturbed by exchanging the partners of a Watson–Crick base pair, that is, by changing a G? C to a C? G or an A? T to a T? A. In contrast, any other combination of bases (e.g., A? G or A ? C) would significantly distort the double helix since the formation of a non-Watson–Crick base pair would require considerable reorientation of the sugar-phosphate chain. B-DNA has two deep exterior grooves that wind between its sugar-phosphate chains as a consequence of the helix axis passing through the approximate center of each base pair. However, the grooves are of unequal size because (1) the top edge of each base pair, is structurally distinct from the bottom edge; and (2) the deoxyribose residues are asymmetric. The **minor groove** exposes that edge of a base pair from which its Clatoms extend whereas the **major groove** exposes the opposite edge of each base pair. Although B-DNA is, by far, the most prevalent form of DNA in the cell, double helical DNAs and RNAs can assume several distinctstructures.



Watson-Crick base pairs. The line joining the Clatoms is the same length in both base pairs and makes equal angles with the glycosidic bonds to the bases. This gives DNA a series of pseudo-twofold symmetry axes (often referred to as dyad axes) that pass through the center of each base pair (red *line*)and are perpendicular to the helix axis. Note that A? T base pairsassociatevia two hydrogen bonds, whereas C? G base pairs are joined bv three hydrogenbonds.

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Chromosome:

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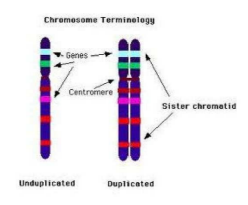
German biologist Walter Flemming in the early 1880s revealed that during cell division the nuclear material organize themselves into visible thread like structures which were named as chromosomes which stains deep with basic dyes. The term chromosome was coined byW.

Waldeyer in 1888. Chrome is coloured and soma is body, hence they mean —colored bodiesl and can be defined as higher order organized arrangement of DNA and proteins. It contains many genes or the hereditary units, regulatory elements and other nucleotide sequences. Chromosomes also contain DNA-bound proteins, which serve in packaging the DNA and control its functions. Chromosomes vary both in number and structure among organisms (Table 1) and the number of chromosomes is characteristic of every species. Benden and Bovery in 1887 reported that the number of chromosomes in each species is constant. W.S. Sutton and T. Boveri in 1902 suggested that chromosomes are the physical structures which acted as messengers ofheredity.

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Chromosomes are tightly coiled DNA around basic histone proteins, which help in the tight packing of DNA. During interphase, the DNA is not tightly coiled into chromosomes,but

exists as chromatin. The structure of a chromosome is given in Figure 2. In eukaryotes to fit the entire length of DNA in the nucleus it undergoes condensation and the degree to which DNA is condensed is expressed as its packing ratio which is the length of DNA divided by the length into which it is packaged into chromatin along with proteins. The shortest human chromosome contains 4.6 x 107 bp of DNA. This is equivalent to 14,000 μ m of extended DNA. In its most condensed state during mitosis, the chromosome is about 2 μ m long. This gives a packing ratio of 7000 (14,000/2). The DNA is packaged stepwise into the higher order chromatin structure and this is known as —hierarchies of chromosomalorganization.



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Chromosome number

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There are normally two copies of each chromosome present in every somatic cell. The number of unique chromosomes (N) in such a cell is known as its haploid number, andthe

total number of chromosomes (2N) is its diploid number. The suffix _ploid' refers to chromosome _sets'. The haploid set of the chromosome is also known as the genome.

Structurally, eukaryotes possess large linear chromosomes unlike prokaryotes which have circular chromosomes. In Eukaryotes other than the nucleus chromosomes are present in mitochondria and chloroplast too. The number of chromosomes in each somatic cell is same for all members of a given species. The organism with lowest number of chromosome is the nematode, *Ascarismegalocephalusunivalens* which has only two chromosomes in the somatic cells(2n=2).

Autosomes and sex chromosomes

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In a diploid cell, there are two of each kind of chromosome (termed homologus chromosomes) except the sex chromosomes. In humans one of the sexhas two of the same kind of sex chromosomes and the other has one of each kind. In humans there are 23 pairs of homologous chromosomes (2n=46). The human female has 44 non sex chromosomes, termed autosomes and one pair of homomorphic sex chromosomes given the designation XX. The human male has 44 autosomes and one pair of heteromorphic sex chromosomes, one X and one Ychromosome.

Morphology:

Size: The size of chromosome is normally measured at mitotic metaphase and may be as short as

 $0.25\mu m$ in fungi and birds to as long as 30 μm in some plants such as Trillium. However, most mitotic chromosome falls in the range of $3\mu m$ in Drosophila to $5\mu m$ in man and $8-12\mu m$ in maize. The monocots contain large sized chromosomes as compared to dicots. Organisms with less number of chromosomes contain comparatively large sized chromosomes. The chromosomes in set vary insize.

Shape: The shape of the chromosome changes from phase to phase in the continuous process of cell growth and cell division. During the resting/interphase stage of the cell, the chromosomes occur in the form of thin, coiled, elastic and contractile, thread like stainable structures, the chromatin threads. In the metaphase and the anaphase, the chromosome becomes thick and filamentous. Each chromosome contains a clear zone, known as centromere or kinetochore,

along their length. The centromere divides the chromosome into two parts and each part is called chromosome arm. The position of centromere varies from chromosome to chromosome providing it a different shape. They could be telocentric (centromere on the proximal end of the chromosome), acrocentric (centromere at one end giving it a very short and another long arm), submetacentric (J or L shaped chromosome with the centromere near the centre), metacentric (v shaped with centromere at thecentre).

Structure of Chromosome

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A chromosome at mitotic metaphase consists of two symmetrical structures called chromatids. Each chromatid contains a single DNA molecule and both chromatidsare attached to each other by centromere and become separated at the beginning of anaphase. The chromomeres are bead like accumulations of chromatin material that are sometimes visible along interphase chromosomes. The chromomere bearing chromatin has an

appearance of a necklace in which several beads occur on a string. Chromomeres are regions

of tightly folded DNA and become especially prominent in polytenechromosomes.

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Centromere in a chromosome contains specific DNA sequences with special proteins bound to them, forming a disc shaped structure, called kinetochore. In electron microscopethe

kinetochore appears as a plate or cup like disc, 0.20-0.25 nm, in diameter situated upon the primary constriction or centromere. The chromosomes of most organisms contain only one centromere and are known as monocentric chromosomes. Some species have diffused centromeres, with microtubules attached along the length of the chromosomes and are termed holocentric chromosomes. Chromosomes of *Ascarismegalocephala*are examples of diffused centromeric chromosomes. Telomere is the chromosomal ends which prevents other chromosomal segments to be fused with it. Besides the primary constrictions or centromeres, chromosomes also possess secondary constrictions are helpful in identifying particular chromosomes in a set. Chromosomes also contain nucleolar organizers which are certain secondary constrictions that contain the genes coding for 5.8S, 18S and 28S ribosomal RNA and induce the formation of nucleoli. Sometimes the chromosomes bear round, elongated or knob like appendages known as satellites. The satellite remains connected with the rest of the chromosomes by a thin chromatin filament

Chromatin

Chemical composition of chromatin

Chromatin consists of DNA, RNA and protein. The protein of chromatin could be of two types:

histones and non-histones.

DNA

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DNA is the most important chemical component of chromatin, since it plays central role of controlling heredity and is most conveniently measured in picograms.

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In addition to describing the genome of an organism by its number of chromosomes, it is also described by the amount of DNA in a haploid cell. This is usually expressed as the amount of

DNA per haploid cell (usually expressed as picograms) or the number of kilobases per haploid cell and is called the C value. This is constant for all cells of a species. For diploid cells it is 2C. Extending the C value we reach the C-value paradox. One immediate feature of eukaryotic organisms highlights a specific anomaly that was detected early in molecular research. Even though eukaryotic organisms appear to have 2-10 times as many genes as prokaryotes, they have many orders of magnitude more DNA in thecell.

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Furthermore, the amount of DNA per genome is correlated not with the presumed evolutionary complexity of a species. This is stated as the **C value paradox**: the amount of

DNA in the haploid cell of an organism is not related to its evolutionary complexity. Lower eukaryotes in generalhave less DNA, such as nematode *Caenorhabditiselegans*which has 20 times more DNA than *E. coli*. Vertebrates have greater DNA content about 3pg, in general about 700 times more than *E. coli*. Salamander *Amphiuma*has a very high DNA content of about 84pg. Man has about 3pg of DNA per haploidgenome.

Histones

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Histones are basic proteins as they are enriched with basic proteins arginine and lysine. At physiological pH they are cationic and can interact with anionic nucleic acids. They form a

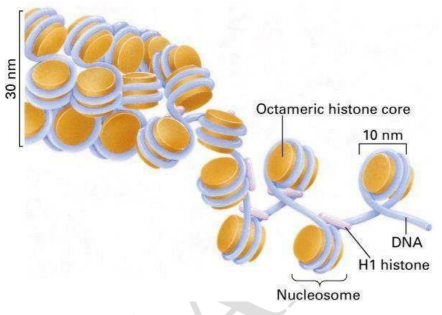
highly condensed structure. The histones are of five types called H1, H2A H2B, H3, and H4which are very similar among different species of eukaryotes and have been highly conserved during evolution. H1 is the least conserved among all and is also loosely bound with DNA. H1 histone is absent in *Sacharomycescerevisiae*.

Non-histones

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In addition to histones the chromatin comprise of many different types of non-histone proteins, which are involved in a range of activities, including DNA replication and gene

expression. They display more diversity or are not conserved. They may also differ between different tissues of same organism. Roger Kornberg in 1974 described the basic structural unit of chromatin which is called thenucleosome.



Euchromatin

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The lightly-stained regions in chromosome when stained with basic dyes are called euchromatin and contain single-copy of genetically-active DNA. The extent of chromatin condensation varies

during the life cycle of the cell and plays an important role in regulating gene expression. In the interphase of cell cycle the chromatin are decondensed and known as euchromatin leading to gene transcription and DNAreplication.

Heterochromatin

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The word heterochromatin was coined by Emil Heitz based on cytological observations. They are highly condensed and ordered areas in nucleosomal arrays. About 10% of interphase chromatin is

called heterochromatin and is in a very highly condensed state that resembles the chromatin of cells undergoing mitosis. They contain a high density of repetitive DNA found at centromeres and telomeres formheterochromatin.

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Heterochromatin are of two types, the constitutive and facultative heterochromatin. The regions that remain condensed throughout the cell cycle are called constitutive heterochromatin whereas

the regions where heterochromatin condensation state can change are known as facultative.

Constitutive heterochromatin is found in the region that flanks the telomeres and centromere of each chromosome and in the distal arm of the Y chromosome in mammals. Constitutive heterochromatin possesses very few genes and they also lead to transcriptional inactivation of nearby genes. This phenomenon of gene silencing is known as —position effect!. Constitutive heterochromatin also inhibits genetic recombination between homologous repetitive sequences circumventing DNA duplications and deletion. Whereas facultative heterochromatin is chromatin that has been specifically inactivated during certain phases of an organism's life or in certain types of differentiated cells. Dosage compensation of X-chromosome or X-chromosome inactivation in mammals is an example of such heterochromatin (Karp 2010). Heterochromatin spreads from a specific nucleation site, causing silencing of most of the X chromosome, thereby regulating genedosage.

Centromeres

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Centromeres are those condensed regions within the chromosome that are responsible for the accurate segregation of the replicated chromosome during mitosis and meiosis. When

chromosomes are stained they typically show a dark-stained region that is the centromere. The actual location where the attachments of spindle fibres occur is called the kinetochore and is composed of both DNA and protein. The DNA sequence within these regions is called *CEN* DNA. Because *CEN* DNA can be moved from one chromosome to another and still provide the chromosome with the ability to segregate, these sequences must not provide any other function. Typically CEN DNA is about 120 base pairs long and consists of several sub-domains, CDE-I, CDE-II and CDE-III. Mutations in the first two sub-domains have no effect upon segregation, but a point mutation in the CDE-III sub-domain completely eliminates the ability of the centromere to function during chromosome segregation. Therefore CDE-III must be actively involved in the binding of the spindle fibers to the centromere. The protein component of the kinetochore is only now being characterized. A complex of three proteins called Cbf-III binds to normal CDE-III regions but cannot bind to a CDE-III region with a point mutation that prevents mitotic segregation. Furthermore, mutants of the genes encoding the Cbf-III proteins also eliminates the ability for chromosomes to segregate during mitosis. Additional analyses of the DNA andprotein

components of the centromere are necessary to fully understand the mechanics of

chromosome segregation.

CDE I	⊂UÆ II	COF II
PUTCACHIG		I CIXIXI GYVIICCGAAyyyyyAAA
ner .	78-86 bp >90% A/T	
10		- 125 bp

The *S. cerevisiae*centrosome. The *S. cerevisae*centromere (CEN) sequences consist of two short conserved sequences (CDE I and CDE III) separated by 78 to 86 base pairs (bp) of ATrich DNA (CDE II). The sequences shown are consensus sequences derived from analysis of the centromere sequences of individual yeast chromosomes. Pu = A or G; x = A or T; y = any base. The figure has been adapted from —The Cell, A Molecular Approachl by Geoffrey M. Cooper, 4th Ed.2007.

Telomeres

Telomeres are the region of DNA at the end of the linear eukaryotic chromosome that are requiredforthereplicationandstabilityofthechromosome.McClintockrecognizedtheirspecial

features when she noticed, that if two chromosomes were broken in a cell, the ends were sticky and end of one could attach to the other and vice versa. However she never observed the attachment of the broken end to the end of an unbroken chromosome suggesting that the end of chromosomes have unique features.

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Telomere sequences remain conserved throughout vertebrates and they form caps that protect the chromosomes from nucleases and other destabilizing influences; and they prevent the ends of

chromosomesfrom fusing with one another. Thetelomeric DNA contains direct tandemly repeated sequences of the form $(T/A)_xG_y$ where x is between 1 and 4 and y is greater than 1. Human telomeres contain the sequence TTAGGG repeated from about 500 to 5000 times. Certain bacteria possess telomeres in their linear genetic material which are of two types; one of the types is called a hairpin telomere. As its name implies, the telomeres bend around from the end of one DNA strand to the end of the complimentary strand. The other type of telomere is known as an invertron telomere. This type acts to allow an overlap between the ends of the complimentary DNAstrands.

Telomere replication

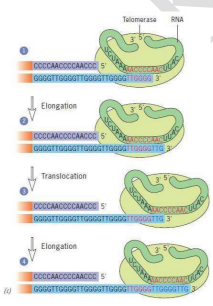
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Telomere replication is an important aspect in DNA replication. The primary difficulty with telomeres is the replication of the lagging strand. Because DNA synthesis requires a RNA template (that provides the free 3'-OH group) to prime DNA replication, and this template is

eventually degraded, a short single-stranded region would be left at the end of the chromosome.

This region would be susceptible to enzymes that degrade single-stranded DNA. The result would be that the length of the chromosome would be shortened after each division. This is known as the end replication problem which is not observed. The action of the telomerase enzymes ensure that the ends of the lagging strands are replicated correctly. Telomerase was discovered in 1984 by Elizabeth Blackburn and Carol Greider of the University of California, Berkeley. It is a reverse transcriptase that synthesizes DNA using an RNA template. Unlike most reverse transcriptases, the enzyme itself contains the RNA that serves as its template, i.e., telomerase can add new repeat units to the 3' end of the overhanging strand. A wellstudied system involves the Tetrahymenaprotozoa organism. The telomeres of this organism end in the sequence 5'-TTGGGG-3'. The telomerase adds a series of 5'-TTGGGGG-3' repeats to the ends of the lagging strand. A hairpin occurs when unusual base pairs between guanine residues in the repeat form. Next the RNA primer is removed, and the 5' end of the lagging strand can be used for DNA synthesis. Ligation occurs between the finished lagging strand and the hairpin. Finally, the hairpin is removed at the 5'-TTGGGG-3' repeat. The replication of telomere has been presented in Figure 4. Telomerase activity is retained in germ cells and zygote and somatic cells after few cell division cycles do not show such activities because otherwise they would divide indefinitely and lead to cancer. Thus telomeres shrink causing chromosome shortening to a critical point when the cell ceases to grow and divide. An inheriteddiseasecalledtheWerner'ssyndromethatcausespatientstoagemuchmorerapidly

abnormal telomere maintenance.



Telomerase replication - Telomerase contains an RNA primer that is complementary to the end of the G-rich strand, which extends past the Crich strand. The telomerase RNA binds to the protruding end of the G-rich strand in step 1 and then serves as a template for the addition of nucleotides onto the 3' terminus of the strand in step 2. After a segment of DNA is synthesized, the telomerase RNA slides to the new end of the strand being elongated in step 3 and serves as the template for the incorporation of additional nucleotides in step 4. The gap in the complementary strand is filled by the replicationenzymes polymerase

 α -primase. This figure has been adapted from Cell and Molecular Biology Concepts and Experiments by Karp,2010.

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Possible Questions

- 1. Describe the experiment of Hershey and Chase on bacteriophages.
- 2. Write in detail about Watson Crick model of DNA.
- 3. Explain in detail about the different types of DNA.
- 4. Write briefly about the Avery experiment on DNA as genetic material.
- 5. Explain in detail about the packing of DNA in histone proteins.
- 6. Write an experiment to prove DNA as genetic material.
- 7. Explain in detail about the Griffith's experiment on Transformation.
- 8. Describe about the super coiling of DNA.
- 9. Write about the phosphodiester bond formation in DNA structure.
- 10. Write about the structural difference between B-form (Watson-Crick) DNA and Z-form DNA.
- 11. Write about Histone proteins.
- 12. What is Euchromatin and Heterochromatin?
- 13. Write short note on chromosomes.
- 14. Write short note on Chargaffs rule..
- 15. Write short note on Okazaki fragments

MOLECULAR BIOLOGY 18BTU302

UNIT II

DNA structure and replication: Replication of DNA in prokaryotes and eukaryotes: Semiconservative nature of DNA replication, Bi-directional replication, DNA polymerases, Replication complex: Pre-priming proteins, primosome, replisome, Rolling circle replication, Unique aspects of eukaryotic chromosome replication, Fidelity of replication.

S.No	HOURS	UNIT II (Total No. of hours - 10 hours)	Page No.
1	1	DNA replication – Introduction	R1 131-135
2	1	Replication of DNA in Prokaryotes	J1 395-420
3	1	Replication of DNA in Eukaryotes	W2
4	1	Semi-conservative nature of DNA replication	T2 109-113
5	1	Bidirectional replication	R1 135-136
6	1	DNA Polymerases	T2 109
7	1	Replication Complexes & Pre-priming proteins	T2 158-162
8	1	Rolling Circle replication	T2 149-151
9	1	Fidelity of replication and Unit Revision	T2 143-144
10	1	Unit test	-

Reference:

T2. Watson, J.D., Baker T.A., Bell, S.P., Gann, A., Levine, M., & Losick, R. (2008).

Molecular Biology of the gene (6th edition). Cold spring Harbour Lab. Press, Pearson Pub.

R1. Lodish et al. (2006). Molecular Cell Biology (5th edition).

W1: www.biologydiscussion.com

DNA replication

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In molecular biology, DNA replication is the biological process of producing two identical replicas of DNA from one original DNA molecule. This process occurs in all living

organisms and is the basis for biological inheritance. DNA is made up of a double helix of two strands, and each strand of the original DNA molecule serves as a template for the production of the complementary strand, a process referred to as semiconservative replication. Cellular proofreading and error-checking mechanisms ensure near perfect fidelity for DNA replication.

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In a cell, DNA replication begins at specific locations, or origins of replication, in the genome. Unwinding of DNA at the origin and synthesis of new strands results inreplication

forks growing bidirectional from the origin. A number of proteins are associated with the

replication fork which helps in terms of the initiation and continuation of DNA synthesis. Most prominently, DNA polymerase synthesizes the new DNA by adding complementary nucleotides to the template strand.

Templated replication

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Sense and antisense strands: The sense strand of a double-stranded DNA molecule has a base sequence similar to that of the RNA that is transcribed from the DNA. The antisense

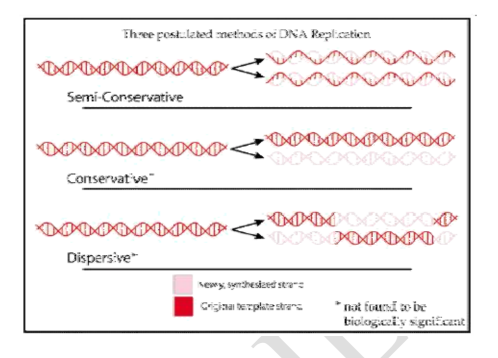
strand (also known as template strand) has a sequence that is the reverse complement of the sense strand (base paired to it with an antiparallel 5' to 3' polarity).

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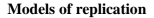
Semiconservative replication: Discovery of the double helical structure of DNA immediately suggested a mechanism for precise replication -- namely separation of the two

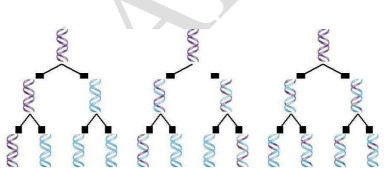
strands and template synthesis of a new copy of each of the missing complementary strands.

This model predicted that each new double helix should have one strand of parental DNA and one strand of newly synthesized DNA.



Three postulated meth ods of replication: 1. Semiconservative, 2. Conservative, 3. Dispersive



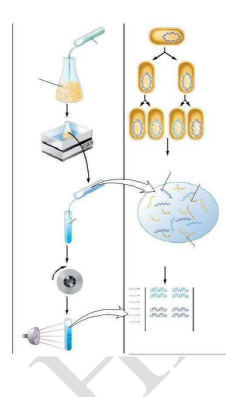


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Messelson-Stahl experiment: Semiconservative replication was demonstrated by starting with bacteria whose DNA was labeled with heavy nitrogen (15N) and then allowing various

amounts of growth to occur in media labeled with normal nitrogen. After one round of replication, the entire DNA had an intermediate density. After two rounds, one half was light and the other half still had an intermediate density. There was no conservation of heavy DNA. However, the data suggested that each of the labeled strands from the original DNA remained intact and separated from its partner in each replication cycle.

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Initiation of replication

• Origin of Replication: The DNA of a bacterial chromosome is a closed circular structure. DNA replication begins at a specific site (origin of replication) characterized by the presence of repeated 9 base and 13 base nucleotide sequences. The repeating units are referred to as 9-mers and 13-mers. The 13 mers are AT rich, making easier to separate the two strands of the double-stranded DNA.

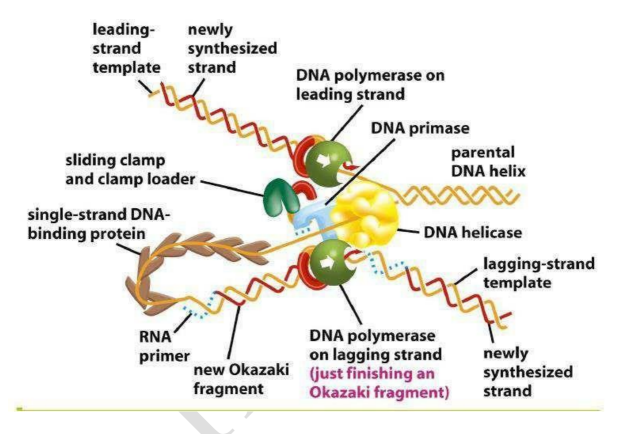
• **DNAa protein:** The protein coded by the DNA a gene binds to the repeated 9mers. This forms a tight loop and generates a strain that causes strand separation in the region containing the AT-rich 13-mers • Helicase: Enzymes called helicases use energy derived from ATP to further separate the two strands of the DNA double helix.• Topoisomerase I: Separation of the two

strands of the DNA double helix requires substantial unwinding of the helix. An enzyme known as topoisomerase I(gyrase) relieves twisting strain that is generated by unwinding the double helix. It is believed to act by cutting one of the strands such that the other strand can rotate freely to relieve the strain, and then resealing the strand that has been temporarily cut.

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• **Keeping the helix open:** Single stranded DNA-binding proteins (SSBPs) attach to the single stranded DNA generated by unwinding the double helix and temporarily keep it from reforming double helical structures.

Patterns of replication



• **Replication forks:** When the two strands of double helical DNA separate and replication of both strands begins, a forked or Y-shaped structure is formed.

• **Bidirectional replication:** In bacterial cells, replication starts at a specific origin of replication within the circular DNA molecule and proceeds in both directions away from the origin. This results in the formation of a replication "bubble", which continues to elongate as replication proceeds. A similar pattern is seen in eukaryotic chromosomes, except that multiple origins are involved.

• Theta structure in bacteria: As replication proceeds in both directions around the circular chromosomes of bacteria, a structure reminiscent of the Greek letter theta is formed.

• **Replicons:** The very long DNA double helices found in eukaryotic chromosomes contain multiple origins of replication, which often initiate bidirectional replication more or less synchronously. Each unit of replication is called a replicon. Replication continues until the replication bubbles fuse to yield fullyreplicated DNA strands.

• Other patterns There are also two alternative strategies for replication of circular DNA that are more complex than the simple bidirectional model. These are the rolling circle mode (sigma mode) used by some types of viruses, and the D loop mode, used for replication of mitochondrial and chloroplast DNAs.

5'-to-3' synthesis

• Unidirectional addition of nucleotides: Polymerization of DNA, and also of RNA, occurs by condensing a 5'-nucleoside triphosphate (dNTP or NTP) onto the 3' hydroxyl group of another nucleotide, or onto the 3'-end of a growing polynucleotide chain. No mechanism exists for extending the 5'-end of a nucleotide chain.

• Energy for synthesis: Hydrolysis of the dNTP or NTP that is being added provides the energy needed to form a covalent bond. The outer two phosphates of the triphosphate are split off and the innermost phosphate forms an ester linkage with the 3'-hydroxyl group at the end of the preexisting chain. This results in a phosphodiester bond

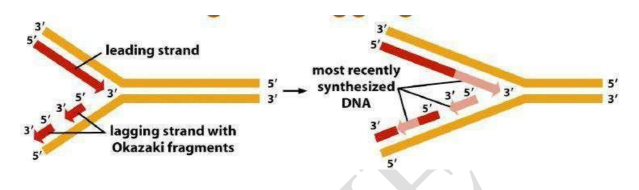
RNA priming

• No new DNA starts: Deoxynucleotides can only be added to 3'-end of a preexisting strand of DNA (or RNA). There are no enzymes capable of initiating the synthesis of a DNA-templated DNA molecule at the level of a single nucleotide. This makes necessary to use an indirect priming procedure.

• **RNA primers:** A variety of enzymes are capable of initiating new DNAtemplated RNA synthesis (as in transcription). New DNA synthesis is primed with a short segment of RNA that is later removed

• **Primase:** A separate enzyme in the initiation complex called primase synthesizes a short RNA primer each time that new DNA synthesis begins, including all new starts in the discontinuous pattern of synthesis described below.

Leading and lagging strands



• Unidirectional synthesis of antiparallel DNA: The inability to synthesize new chains in a 3' to 5' direction adds a major complication to the replication of antiparallel double-stranded DNA. At any replication fork, one of the template strands has a 3' to 5' orientation, which is what is needed for synthesis of a new complementary strand in a 5' to 3' direction. Synthesis on that template is primed and starts very quickly. However, the other template strand has a 5' to 3' orientation and is thus unable to support synthesis beginning at the origin and moving away from it in a 3' to 5' direction. Because of this, the 5' to 3' template strand accumulates in a single stranded configuration until there is a sufficient length so that synthesis of its complementary antiparallel strand can be primed and initiated in a 5'-to-3' direction ("backward" toward the origin of replication). The strand whose synthesis begins immediately is called the "leading" strand, and the one whose synthesis is delayed is called the "lagging" strand.

• **Discontinuous synthesis -- Okazaki fragments:** As synthesis of the leading strand continues, more and more of the single stranded DNA of the lagging strand is unwound. Each time that a sufficient length is reached, synthesis of a new segment of the complementary strand begins. If the replicating DNA is denatured (separated into individual strands) before the newly synthesized pieces of the lagging strand have been ligated together, a number of relatively small fragments of newly synthesized DNA will be recovered, together with the much longer strands

produced by continuous synthesis in the leading strand. The small fragments are called "Okazaki fragments", named for the person who first discovered them.

DNA polymerase III

Template DNA synthesis: After the RNA primer has reached an adequate length, DNA polymerase III begins synthesis of DNA, which proceeds to completion in the leading strand, and proceeds until the 5'-end of the previous primer is encountered in the lagging strand.

• **DNA polymerase III holoenzyme:** DNA polymerase III is a highly complex dimeric aggregate, consisting of 20 or more protein subunits. The alpha subunits perform the actual DNA synthesis, but operate in conjunction with multiple accessory proteins

• Simultaneous synthesis of leading and lagging strands: There is yet another complicating factor in DNA synthesis. It is now generally believed that the leading and lagging strands are synthesized simultaneously by a single dimeric DNA polymerase III complex. This requires formation of a looped structure with the leading and lagging strands so positioned that their synthesis can occur side by side in the same orientation, despite the fact that the newly synthesized chains are growing in opposite directions relative to the overall DNA that is being replicated. A similar looping also appears to occur in eukaryotic DNA replication.

• **Clamping function of beta subunits:** The beta subunits appear to have a clamping function that keeps the leading and lagging strands appropriately aligned with the catalytically active alpha subunits.

• **Replisome complex:** The entire DNA-synthesizing complex at each replication fork, which also includes topoisomerase, helicase, and primase, is sometimes referred to as a replisome.

DNA polymerase I

• Not the primary enzyme for DNA synthesis: DNA polymerase I is so named because it was the first of the DNA polymerase enzymes to be isolated and characterized. Although it is capable of template-directed DNA synthesis, it is now known not to be the enzyme primarily responsible for new DNA synthesis, but it does have other important roles, as described below.

• 5' to 3' exonuclease activity: DNA polymerase I has an unusual 5' to 3' exonuclease activity. This gives it the ability to start at a single-stranded break and progressively remove nucleotides and replace them in a 5' to 3' direction.

• **Removal of RNA primer:** In the lagging strand, when DNA polymerase III runs into the 5'end of the previous RNA primer, it is unable to proceed further. Although the new DNA butts up against the primer, it is not covalently joined to it. DNA polymerase III dissociates, leaving a "nick" (a single stranded gap) between the new DNA and the primer. Starting at the nick, DNA polymerase I removes the primer ribonucleotides one at a time, using its 5' to 3' exonuclease activity, and replaces them with deoxyribonucleotides, using its DNA polymerase activity..

• **DNA repair:** DNA polymerase I is also used to fill in short gaps in DNA, often as a part of a repair process that excises part of a damaged strand and replaces it with new DNA templated from the remaining strand. Proofreading and DNA repair

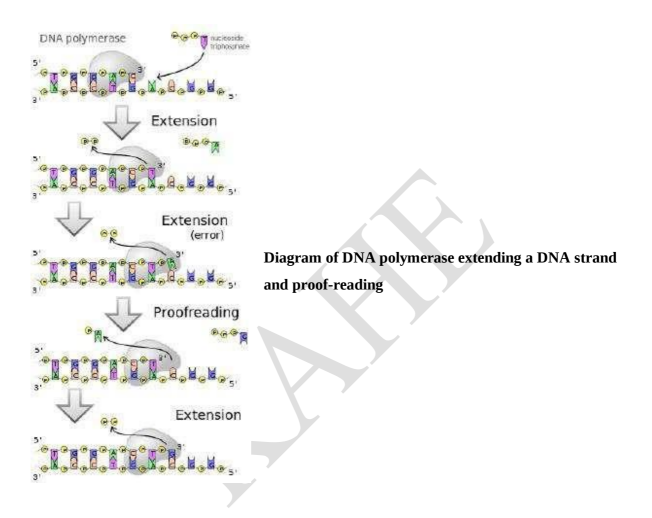
• Removal of mismatched bases: DNA polymerases III and I both have 3' to 5' exonuclease activity. This allows them to remove a mismatched nucleotide that has just been added to a growing DNA chain and make another attempt to insert the correct nucleotide. This "proofreading" function helps to reduce the number of mistakes in DNA synthesis that would otherwise result in mutation

•DNA polymerase II: There is yet another prokaryotic DNA polymerase, designated DNA polymerase II, whose main function appears to be to synthesize replacement DNA during DNA repair.

Final steps in DNA synthesis

• DNA Ligase: After the last ribonucleotide is removed from the primer and replaced with a deoxynucleotide, there is still a nick in the newly synthesized lagging strand. This nick is closed by DNA ligase, which forms a covalent phosphodiester bond between the Okazaki fragments, joining them into a continuous strand of DNA. Note that Okazaki fragments accumulate when ligase function is impaired.

• Topoisomerase II: Its role is to cut and reseal the newly synthesized DNAs as needed so that they can separate from each other (it is easy to visualize two circular genomes linked through each other at the end of replication).



Rolling cycle method of Plasmid DNA

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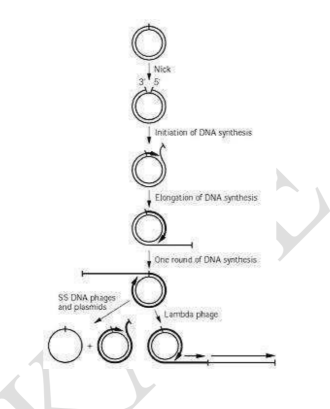
In DNA replication, the DNA polymerase cannot initiate the synthesis of a new DNA strand and must rely on a priming device. In general, an RNA primer is synthesized at or near a

replication origin to start synthesis of the leading strand. However, a DNA primer terminus can be generated by a nuclease-generated nick at a specific place in some circular duplex DNA, and replication will then proceed unidirectionally. This mode of replication is called rolling circle replication and is found for replication of the replicative form (RF) form of

bacteriophage single-stranded genomes of Gram-negative bacteria and of the multicopy plasmids of Gram-positive bacteria (see Single-Stranded DNA Replication).

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Rolling circle replication is also observed in the late stage of the replication of the lambda phage genome and in the process of the conjugative transfer of bacterial plasmids.



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DNA synthesis initiates using the free 3'-OH end at the nick as a primer, and a replication fork proceeds around the template. In the process, the newly synthesized strand displaces the

old strand from the template.

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In the case of replication of the RF form of single-stranded phage genomes and of plasmids of Gram-positive bacteria, the displaced old strand is cleaved off after one round of replication and is converted into the circular, double-stranded form. In contrast, in phage lambda replication, the replication fork proceeds a number of revolutions around the template without cleavage of the displaced strand, and the displaced strand becomes double-stranded

as it is peeled off.

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The linear concatemer thus created is cleaved into one unit length and packaged into the phage particles. In the conjugation process of plasmids, the displaced strand is transferred.

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The initial nick of rolling circle replication is introduced by an endonuclease specific for each system. The similarity of the amino acid sequences of the initiator endonucleases and of the

proteins (known as relaxases) involved in the initiation and termination of conjugative DNA strand transfer clearly indicate that they are evolved from a common ancestor.

Rolling cycle method of DNA replication in ϕ X 174

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All circular genomes do not follow the same pattern of replication described in *E. coli*. In some bacteriophages (phi, lambda and phi X 174), in mitochondrial chromosomes and during

bacterial mating an alternative method known as the rolling circle has been demonstrated.

This method as seen in the small bacteriophage ϕ X 174 will be described here.

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 ϕ X174 has a single stranded DNA molecule only about 6,000 nucleotides in length. Its newly formed DNA is linear, not circular. The process takes place in following way. Soon

after φ X174 enters the host E. coli cell its single stranded parent DNA (designated + strand) synthesizes a complementary minus strand (with the help of host cell enzymes) to form duplex DNA (+ -). This is the replicative form (RF) of φ XI74.

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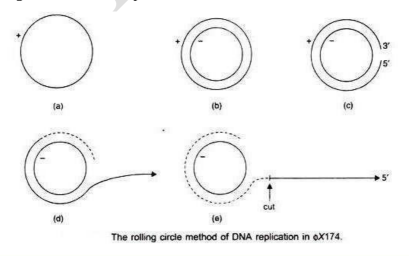
An endonuclease enzyme produces a nick in the + strand, exposing a free 3' end and a free 5' end. DNA synthesis begins by addition of deoxyribonucleotides by the enzyme DNA

polymerase to the free 3' end of the + strand using the strand as template.

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The minus strand revolves while serving as template hence the name rolling circle. The addition of nucleotides displaces the free 5' end outwards in the form of a free tail. The

rolling circle revolves a number of times thus increasing the length of the + strand by a corresponding number of + complements.



It appears that the tail is finally cut into the correct genome lengths by an endonuclease to provide as many copies of free, linear DNA molecules. The free ends joined by the newly formed linear DNA are joined by the enzyme DNA ligase to form closed, circular progeny chromosomes.

Fidelity of Replication

Since a single polypeptide as small as the Pol I Klenow fragment can replicate DNA by itself, why does *E. coli* maintain a battery of >20 intricately coordinated proteins to

replicate its chromosome? The answer apparently is to ensure the nearly perfect fidelity of DNA replication required to preserve the genetic message's integrity from generation to generation. The rates of reversion of mutant E. coli or T4 phage to the wild type indicates that only one mispairing occurs per 10^8 to 10^{10} base pairs replicated. This

corresponds to _1error per 1000 bacteria per generation. Such high replication accuracy arises from four sources:

1. Cells maintain balanced levels of dNTPs through the mechanism discussed in Section 28-3Ad. This is an important aspect of replication fidelity because a dNTP present at aberrantly high levels is more likely to be misincorporated and, conversely, one present at low levels is more likely to be replaced by the dNTPs present at higher levels.

2. The polymerase reaction itself has extraordinary fidelity. This is because, as we have seen (Section 30-2Ae), the polymerase reaction occurs in two stages: (1) a binding step in which the incoming dNTP base-pairs with the template while the enzyme is in an open conformation that cannot catalyze the polymerase reaction; and (2) a catalysis step in which the polymerase forms a closed conformation about the newly formed base pair, which properly positions its catalytic residues (induced fit). Since the formation of the closed conformation requires that the incoming dNTP form a Watson–Crick-shaped base pair with the template, the conformation change constitutes a double check for correct base pairing.

3. The 3_i , S 5_i exonuclease functions of Pol I and Pol III detect and eliminate the occasional errors made by their polymerase functions. In fact, mutations that increase a DNA polymerase's proofreading exonuclease activity decrease the rates of mutation of other genes.

4. A remarkable battery of enzyme systems, contained in all cells, function to repair residual errors in the newly synthesized DNA as well as any damage that it may incur after its synthesis through chemical and/or physical insults.

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In addition, the inability of a DNA polymerase to initiate chain elongation without a primer is a feature that increases DNA replication fidelity. The first few nucleotides of a

chain to be coupled together are those most likely to be mispaired because of the cooperative nature of base pairing interactions. The editing of a short duplex oligonucleotide is similarly an error-prone process.

The use of RNA primers eliminates this source RNA is eventually replaced by DNA under conditions that permit accurate base pairing to be achieved. One might wonder why cells have evolved the complex system of discontinuous lagging strand synthesis rather than a DNA polymerase that could simply extend DNA chains in their 3'- 5' direction. Consideration of the chemistry of DNA chain extension also leads to the conclusion that this system promotes high-fidelity replication. The linking of 5'-deoxynucleotide triphosphates in the 3'- 5' direction would require the retention of the growing chain's 5'- terminal triphosphate group to drive the next coupling step.

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On editing a mispaired 5'-terminal nucleotide, this putative polymerase would—in analogy with Pol I, for example excise the offending nucleotide, leaving either a 5'-OH

or a 5'-phosphate group. Neither of these terminal groups is capable of energizing further chain extension. A proofreading 3'- 5' DNA polymerase would therefore have to be capable of reactivating its edited product. The inherent complexity of such a system has presumably selected against its evolution.

KARPAGAM ACADEMY OF HIGHER EDUCATION				
CLASS: II BSC BT	COURSE	NAME: MOLECULAR BIOLOGY		
COURSE CODE: 18BTU302	UNIT: II	BATCH-2018-2021		

Possible Questions

- 1. Explain in detail on semiconservative nature of DNA replication
- 2. Elaborate on various steps involved in DNA replication in eukaryotes.
- 3. Explain about the bidirectional property of DNA replication.
- 4. Write in detail about the role of different polymerases enzyme in replication.
- 5. Explain the fidelity of replication
- 6. Describe the experiment of Meselson and stahl on replication.
- 7. Write briefly on Rolling circle replication.
- 8. Describe the various steps involved in eukaryotic replication.
- 9. Explain in detail about the semi-discontinuous property of DNA.
- 10. Describe the formation of replication fork.

KARPAGAM ACADEMY OF HIGHER EDUCATION

CLASS: II BSC BTCOURSE NAME: MOLECULAR BIOLOGYCOURSE CODE: 18BTU302UNIT: IIIBATCH-2018-2021

MOLECULAR BIOLOGY 18BTU302

UNIT III

Transcription and RNA processing: RNA structure and types of RNA, Transcription in prokaryotes: Prokaryotic RNA polymerase, role of sigma factor, promoter, Initiation, elongation and termination of RNA chains Transcription in eukaryotes: Eukaryotic RNA polymerases, transcription factors, promoters, enhancers, mechanism of transcription initiation, promoter clearance and elongation RNA splicing and processing: processing of pre-mRNA: 5' cap formation, polyadenylation, splicing, rRNA and tRNA splicing.

UNIT III (Total No. of hours - 11 hours)				
18	1	RNA structure and types of RNA	R2 60-64	
19	1	Transcription in Prokaryotes	T2 361-363	
20	1	Transcription in Eukaryotes	T2 363-366	
21	1	RNA Polymerases and role of σ factor		
22	1	Promotor, Initiation, Elongation and termination	D2 215 262	
23	1	Eukaryotic RNA Polymerses		
24	1	Transcription factors		
25	1	Mechanism of transcription Initiation		
26	1	RNA splicing and Processing	T2 379	
27	1	Processing of different RNA's	T2 412-415	
28	1	Unit test	-	

- T2. Watson, J.D., Baker T.A., Bell, S.P., Gann, A., Levine, M., & Losick, R. (2008). Molecular Biology of the gene (6th edition). Cold spring Harbour Lab. Press, Pearson Pub.
- R2. David Friefielder (2006). Molecular Biology (2006). Narosa Publishing house.

RNA

 \triangleright

RNA differs from DNA in both structural and functional respects. RNA has two major structural differences: each of the ribose rings contains a 2'-hydroxyl, and RNA uses uracil in

place of thymine.

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RNA molecules are capable of base pairing, but generally will not form large regions of stable RNA-RNA double helix.

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RNA can act as a genetic material (although this role, at least for current organisms, seems to be restricted to viruses). Unlike DNA, RNA can form complex three-dimensional structures.

As a result, RNA can also exhibit catalytic activity.

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The combination of the ability to store genetic information with the ability to catalyze reactions has resulted in a proposal for the origin of life: the "RNA World".

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The RNA world hypothesis proposes that RNA molecules once filled all of the roles of protein and nucleic acid macromolecules, and acted in both an information storage capacity

and as the source of the enzymatic activity required for metabolic reactions. In general, RNA

is less suited to acting as genetic material than DNA, and is less suited to forming efficient

catalysts than proteins.

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Assuming that the RNA world once existed, nearly all of its functions have been taken over by other biological molecules. However, some vestiges of the RNA world may still exist.

The vast majority of RNA functions are concerned with protein synthesis.

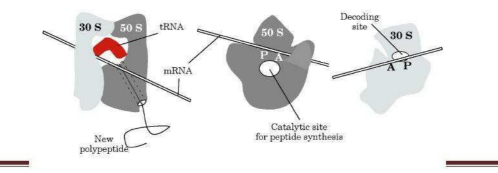
The major types of RNA

Ribosomal RNA (rRNA)

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Ribosomal RNA molecules comprise 65 to 70% of the mass of the ribosome (the machinery responsible for protein synthesis). Ribosomes are very large objects; prokaryotic ribosomes

have molecular weights of about 2.5 million, while eukaryotic ribosomes have molecular weights of about 4 million.



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The original studies on ribosomes used relatively crude techniques that were unable to measure size in terms of molecular weight. Instead the size of the ribosomal particles and

their components were measured by their rate of sedimentation (movement driven by gravitational acceleration or centrifugal acceleration).

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Sedimentation is a function of size, shape, and density, with larger objects tending to sediment faster than smaller ones. Object sizes are measured in Svedberg units. Prokaryotic

ribosomes are 70 S particles, with each comprised of a large (50 S) and a small (30 S) subunit. Eukaryotic ribosomes are 80 S particles, comprised of a large (60 S) and a small (40

S) subunit. You will notice that the Svedberg units are not additive for the particles sizes; this is due to the effects of shape on sedimentation.

\triangleright

The eukaryotic 40S Ribosome contains 1 rRNA (18 S rRNA = 1900 bases) and about 35 different proteins. The 60S ribosome contains 3 rRNA (5 S = 120 bases, 5.8 S = 160 bases,

and 28 S = 4700 bases), and about 50 proteins. The 5 S rRNA has its own gene; the others are synthesized as a single transcript that is then cleaved to release the mature RNA molecules that become part of the ribosome. Until relatively recently, it was assumed that the ribosomal RNA performed a largely structural function.

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However, more recent data strongly suggests that the rRNA acts as the enzyme, with the protein acting as the structural scaffolding. These data include results from the recent high-

resolution (2.4 Å) X-ray diffraction structure of the large subunit and low-resolution (5 Å) structure of the complete ribosome from the bacterium *Haloarcula marismortui*. Examination of the high-resolution structure of the large subunit, and of the lower resolution structures of the entire particle (used to generate the cartoons above) strongly suggests that only RNA is present at the catalytic site.

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In the structure below, proteins are shown in black, and the orientation of the large subunit is similar to the center cartoon above. Examination of this structure suggests that no protein is

present in the catalytic site.

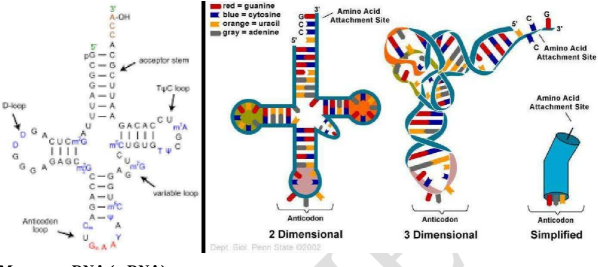
Transfer RNA (tRNA)

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tRNA is a ~75 base molecule that carries the amino acids, and transfers them to the growing protein. tRNAs are thought to have a common tertiary structure structure based

on X-ray diffraction analysis is shown below. Analysis of the tRNA sequence suggests a

cloverleaf secondary structure formed by regions of base pairing between the sections of the RNA strand, with this cloverleaf folding into the three-dimensional structure.



Messenger RNA (mRNA)

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mRNA molecules contain the coding sequence for proteins. The mRNA molecules can vary considerably in size, with eukaryotic transcripts including the largest known

ribonucleic acids. This is most obvious before splicing of introns, because many transcripts exceed 100 kb in length.

snRNA small nuclear ribonucleic acid (snRNA),

\triangleright

It is also commonly referred to as **U-RNA**, is a class of small RNA molecules that are found within the splicing speckles and Cajal bodies of the cell nucleus in eukaryotic cells. The

length of an average snRNA is approximately 150 nucleotides. They are transcribed by either RNA polymerase II or RNA polymerase III, and studies have shown that their primary function is in the processing of pre- messenger RNA (hnRNA) in the nucleus. They have also been shown to aid in the regulation of transcription factors (7SK RNA) or RNA polymerase II (B2 RNA), and maintaining thetelomeres.

 \triangleright

snRNA are always associated with a set of specific proteins, and the complexes are referred to as small nuclear ribonucleoproteins (snRNP, often pronounced "snurps"). Each snRNP

particle is composed of several Sm proteins, the snRNA component, and snRNP-specific proteins. The most common snRNA components of these complexes are known, respectively, as: U1 spliceosomal RNA, U2 spliceosomal RNA, U4 spliceosomal RNA, U5 spliceosomal

RNA, and U6 spliceosomal RNA. Their nomenclature derives from their high uridine content.

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snRNAs were discovered by accident during a gel electrophoresis experiment in 1966. An unexpected type of RNA was found in the gel and investigated. Later analysis has shown that

these RNA were high in uridylate and were established in the nucleus.

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A large group of snRNAs are known as small nucleolar RNAs (snoRNAs). These are small RNA molecules that play an essential role in RNA biogenesis and guide chemical

modifications of ribosomal RNAs (rRNAs) and other RNA genes (tRNA and snRNAs). They are located in the nucleolus and the Cajal bodies of eukaryotic cells (the major sites of RNA synthesis), where they are called scaRNAs (small Cajal body-specific RNAs).

Classes of sn RNA

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snRNA are often divided into two classes based upon both common sequence features as well as associated protein factors such as the RNA-binding LSm proteins.

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The first class, known as **Sm-class snRNA**, is more widely studied and consists of U1, U2, U4, U4atac, U5, U7, U11, and U12. Sm-class snRNA are transcribed by RNA polymerase II.

The pre-snRNA are transcribed and receive the usual 7-methylguanosine five-prime cap in the nucleus. They are then exported to the cytoplasm through nuclear pores for further processing. In the cytoplasm, the snRNA receive 3' trimming to form a 3' stem-loop structure, as well as hypermethylation of the 5' cap to form trimethylguanosine. The 3' stem structure is necessary for recognition by the survival of motor neuron(SMN) protein. This complex assembles the snRNA into stable ribonucleoproteins (RNPs). The modified 5' cap is then required to import the snRNP back into the nucleus. All of these uridine-rich snRNA, with the exception of U7, form the core of the spliceosome. Splicing, or the removal of introns, is a major aspect of post-transcriptional modification, and takes place only in the nucleus of eukaryotes. U7 snRNA has been found to function in histone pre-mRNA processing.

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The second class, known as **Lsm-class snRNA**, consists of U6 and U6atac. Lsm-class snRNAs are transcribed by RNA polymerase III and never leave the nucleus, in contrast to

Sm-class snRNA. Lsm-class snRNAs contain a 5'-\gamma-monomethylphosphate cap and a 3'

stem–loop, terminating in a stretch of uridines that form the binding site for a distinct heteroheptameric ring of Lsm proteins.

SnRNA in the splicosome

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Spliceosomes are a major component of an integral step in eukaryotic precursor messenger RNA maturation. A mistake in even a single nucleotide can be devastating to the cell, and a

reliable, repeatable method of RNA processing is necessary to ensure cell survival. The spliceosome is a large, protein-RNA complex that consists of five small nuclear RNAs (U1, U2, U4, U5, and U6) and over 150 proteins.

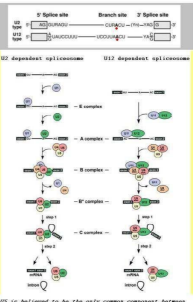
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The snRNAs, along with their associated proteins, form ribonucleoprotein complexes (snRNPs), which bind to specific sequences on the pre-mRNA substrate. This intricate

process results in two sequential transesterification reactions. These reactions will produce a free lariat intron and ligate two exons to form a mature mRNA. There are two separate classes of spliceosomes. The major class, which is far more abundant in eukaryotic cells, splices primarily U2-type introns.

The initial step of splicing is the bonding of the U1 snRNP and its associated proteins to the 5' splice end to

the hnRNA. This creates the commitment complex which will constrain the hnRNA to the splicing pathway. Then, U2 snRNP is recruited to the spliceosome binding site and forms complex A. U2 snRNP changes the conformation of the hnRNA-snRNP complex, exposing the nucleotide favorably for splicing. Following the conformation change, the U4/U5/U6 tri-snRNP complex binds to complex A to form the structure known as complex B. After rearrangement, complex C is formed, and the spliceosome is active for catalysis.



US is believed to be the only common component betwee major and minor spliceosomes. Reference: Vill CL, Luhrmann R.Biol Chem. 2005 Aug:386(8):713–24.

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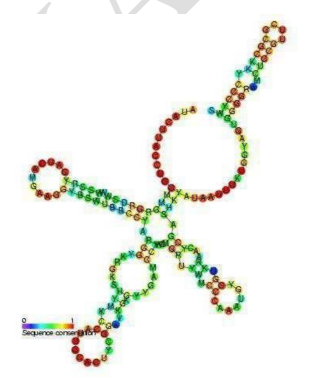
In addition to this main spliceosome complex, there exists a much less common (~1%) minor spliceosome. This complex comprises U11, U12, U4atac, U6atac and U5 snRNPs. These

snRNPs are functional analogs of the snRNPs used in the major spliceosome. The minor spliceosome splices U-12 type introns. The two types of introns mainly differ in their

splicing sites: U2-type introns have GT-AG 5' and 3' splice sites while U12-type introns have AT-AC at their 5' and 3' ends. The minor spliceosome carries out its function through a different pathway from the major spliceosome.

U1 snRNA

U1 snRNP is the initiator of spliceosomal activity in the cell by base pairing with the hnRNA. In the major spliceosome, experimental data has shown that the U1 snRNP is present in equal stoichiometry with U2, U4, U5, and U6 snRNP. However, U1 snRNP's abundance in human cells is far greater than that of the other snRNPs. Through U1 snRNA gene knockdown in HeLacells, studies have shown the U1 snRNA holds great importance for cellular function. When U1 snRNA genes were knocked out, genomic microarrays showed an increased accumulation of unspliced pre-mRNA. In addition, the knockout was shown to cause premature cleavage andpolyadenylation primarily in introns located near the beginning of the transcript. When other uridine based snRNAs were knocked out, this effect was not seen. Thus, U1 snRNA-pre-mRNA base pairing was shown to protect pre-mRNA from polyadenylation as well as premature cleavage. This special protection may explain the overabundance of U1 snRNA in the cell.



Transcription

• Transcription is a fundamental cellular process: RNA polymerases "transcribe" the genetic information on DNA into RNA strands. All cells have RNA polymerases (RNAP).

• The RNA polymerases increase in complexity as you go from viruses (example, T7

RNA polymerase is made up of a single protein), to bacterial systems (one RNA polymerase made up of the proteins - beta, beta', 2 x alpha, omega and the sigma factor), and finally to eukaryotic systems (Three RNA polymerases - Pol I, Pol II, and Pol III, each with ten or more subunits).

• While the RNA polymerases have become increasing complex as life evolved, their overall structure (as evidenced by crystallographic structures of bacterial RNA polymerase and Pol II) show remarkable similarity. There is also sequence similarity between the bacterial polymerase protein subunits and the proteins that make up the eukaryotic polymerases.

Transcription of any gene usually involves three distinct stages:

1. First, the RNA polymerase has to find the start site of a gene. The "holoenzyme" form of the polymerase does this by looking for the "promoter" site that exists just upstream of the gene start site. This process is termed "transcription initiation". This is followed by opening up (melting) of the duplex DNA to form an "open complex".

2. This is followed by a rapid change into the "elongation" phase of transcription where the "core polymerase" part of the RNA polymerase rapidly transcribes an RNA strand that is complementary to the "template" strand of the DNA. The change into the elongation phase usually occurs after a few bases of RNA have been transcribed (typically about 8-9 bases of RNA in bacterial system which form a RNA-DNA hybrid with the template strand), and involves a "clamping down" on the DNA to prevent the polymerase from falling off the DNA.

3. The final stage of the transcription of a gene is "termination", after the stop codon of the gene. The process of termination usually involves sequences where the polymerase slows down or stalls, and the polymerase-RNA-DNA complex (often proteins such as rho and NusA are involved in bacterial systems).

• Genes have to be transcribed to mRNAs before they can be translated into proteins; more or less mRNA from a particular gene equals more or less of the protein encoded by the gene.

Transcription is, thus, an important point in the control of gene "expression". Most genes are controlled transcriptionally, usually by regulation of the level of transcriptional initiation. For example, if a gene has a strong promoter, it will be more highly expressed when compared to another gene with a weak promoter site. Similarly if a regulatory protein can bind the promoter site of a gene (and prevent transcription initiation), then it can turn off the expression of that gene.

• Transcriptional control of genetic expression is vital for cellular functions, and many diseases and cancers are results of defects in the transcriptional control of essentials genes.

Transcription initiation

• Transcription initiation in bacteria (prokaryotes) involves sigma factors. The sigma factor combine with the core RNA polymerase to form a holoenzyme that is competent for promoter binding. The core RNA polymerase, by itself, cannot bind the promoter site.

• The sigma factor can be thought of as the specificity factor in the RNA polymerase.

Each bacteria has several different sigma factors that recognize slightly different promoter sequences. Predominant among these is the sigma70 (70 kDa in E. coli; also called rpoD), which initiates the transcription of most genes in exponentially growing cells. There are two general classes of sigma factors - sigma 70 class and sigma 54 class:

• The sigma 70 class of sigma factors share extensive sequence homology, and bind to two conserved sequences upstream of the gene start site (the -35 box and the -10 box). Each of these sigma factors recognizes slight variations in these conserved sequence boxes. Sigma70 binds promoter DNA as a part of the holoenzyme, and binds DNA very poorly in absence of the core enzyme.

• The sigma54 class of genes (54 kDa in E. coli; also called rpoN) controls a much smaller set of genes than sigma70. It recognizes different conserved sequences (the -12 and -24 boxes). Unlike Sigma70, sigma54 can bind DNA even in the absence of the core RNA polymerase. It however lacks the ability to melt promoter DNA on its own - for this it needs to interact with other activator proteins that bind further upstream of the promoter site, as well as the core RNAP.

• In eukaryotic systems, transcription initiation is very different. There are no sigma factors. Instead, the central protein in forming the "pre-initiation complex " (PIC) is the TATA binding protein (TBP), that binds to the TATA box, a conserved sequence just upstream of the

initiation region. A large number of other general transcription factors such as TFIIB, TFIIE, TFIIF, TFIIH (TF stands for transcription factor; II stands for Pol II; there are similar factors for Pol I and Pol II) and others assemble to form the multisubunit TFIID complex. This PIC then recruits the RNA polymerase (Pol I, II, or III in eukaryotic cells) to initiate transcription. The PIC often remains at the promoter site, and is then available to initiate another round of transcription.

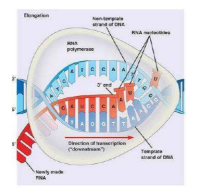
• TBP is a universal transcription factor, and is seen in all eukaryotes and archaea. It sharply bends DNA at the TATA box.

Transcription elongation

• Once initiation is complete and the open complex forms, the RNA polymerase begins to read the template strand and add corresponding RNA nucleotides. This process is not always efficient, and the polymerase may make several passes at this. After a long enough RNA-DNA hybrid is made, the polymerase clears the promoter region and moves rapidly downstream. This is preceeded by a large conformational change in the polymerase core enzyme, as it clamps down on the DNA and becomes quite processive.

• While transcription elongation is quite rapid, the polymerase does not transcribe all sequences with equal efficieny. The elongation rate is not uniform, and there can be pausing or stalling. Elongation factors (GreA/GreB in bacterial systems; TFIIS in eukaryotes) act to help the polymerase along by stimulating backtracking and cleavage of the newly formed RNA (from the 3' end).

• Other factors are involved in the elongation cycle - for example, in eukaryotic Pol II there are the elongins and ELL proteins that increase the elongation rate, as well as factors to remodel chromatin.



Transcription elongation

Transcription activation & repression

• There are multiple modes of transcription activation. All these usually involve different protein factors (activators) that bind DNA sequences (enhancer sequences) in and around the promoter site. All these act to increase the affinity of the initiation complex or RNA holoenzyme at the promoter (thus enhancing the chance that a productive open complex will form and transcription will initiate).

• When enhancer elements combine with poor promoter sequences, activators can modulate the activity of a gene by several hundred or thousand fold. For example, if a gene has a poor -35 or -10 box in its promoter site, it will be poorly expressed - an activator protein can, in such a case, enhance the activity of the gene by several orders of magnitude, by recruiting the RNA polymerase (or other initiation factors) to the promoter site.

• Repression, on the other hand, works by having protein (repressors) that sit on or close to the promoter regions of the DNA, preventing RNA polymerase or initiator/activator proteins from starting transcription initiation (thereby "turning-off" the gene).

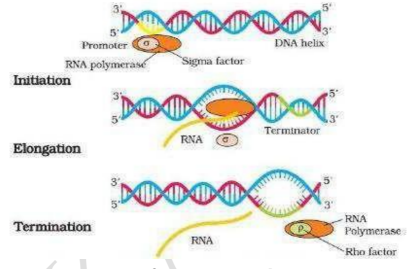
• Activators and repressors are usually DNA binding proteins. These proteins have common DNA binding motifs such as Zn-fingers, helix-turn-helix, etc. Activators also have regions that interact with different domains of the RNA polymerase (parts of alpha, sigma, etc.).

• In addition to protein activators, there are also DNA sequences that can directly interact with RNA polymerase components. For example, the alpha subunit in the bacterial polymerase has two domains - the alpha NTD (n-terminal domain) and the alpha CTD (c-terminal domain), linked by a flexible linker region. The alpha CTD can bind DNA sequences (UP elements) upstream of the promoter region, enhancing the affinity of the polymerase on the promoter.

Transcription termination

• Termination in bacterial system can be broadly classified as "rho independent", and "rho dependent". o In rho independent termination, there is formation of a stable GC rich stemloop in the newly synthesized RNA followed by a string of U's (A's in the template strand) spaced about 20 bases downstream (these sites are often called intrinsic terminators). The stem loop "snares" the polymerase, slowing or stalling it. This pause, coupled with the low stability of the RNA-DNA hybrid at the active site (run of A=U basepairs) allows the RNA polymerase to fall off the template DNA and terminates the RNA transcription for that gene. o In rho dependent termination, rho binds the newly formed RNA (as hexamers), and stalls the RNA polymerase by interacting with it. In some models, the Rho hexamer translocates on the RNA. Rho termination activity is stimulated by ATP hydrolysis. This activity is greatly enhanced by Nus factors.

• In eukaryotic cells, transcription termination involves cleavage of the elongating RNA chain by specific endonucleases which recognize particular sequences (AAUAAA) in the newly formed RNA. Once this happens, the RNA elongation complex is destabilized, and falls off the DNA. It is then available to attach to another nearby PIC, and start transcribing again.



Process of transcription in bacteria

RNA processing (for eukaryotes)

Eukaryotic genes contain Exons (which encode information that will end up being translated into protein) separated by Introns (sequences that will not encode proteins). At the 5' end of a gene we find a promoter and often a CpG island. Both of these elements regulate the transcription of a gene. At the 3' end of a gene we find a stop sequence, and a signal for polyadenylation (AAUAAA). In Eukaryotes this primary RNA transcript is then processed into a messenger RNA (mRNA) if the gene is to be translated into protein. This involves removing introns, which do not encode for protein, and splicing the remaining exons together. At the 5' end of the mRNA a 7-Methyguanosine residue is added to provide a protective cap. At the 3' end of the mRNA, 50-100 adenosine residues are added, generating a poly A tail.

REVERSE TRANSCRIPTION

Some viruses (such as HIV, the cause of AIDS), have the ability to transcribe RNA into DNA. HIV has an RNA genome that is duplicated into DNA. The resulting DNA can be merged with the DNA genome of the host cell. The main enzyme responsible for synthesis of DNA from an RNA template is called reverse transcriptase. In the case of HIV, reverse transcriptase is responsible for synthesizing a complementary DNA strand (cDNA) to the viral RNA genome. An associated enzyme, ribonuclease H, digests the RNA strand, and reverse transcriptase synthesizes a complementary strand of DNA to form a double helix DNA structure. This cDNA is integrated into the host cell's genome via another enzyme (integrase) causing the host cell to generate viral proteins that reassemble into new viral particles. In HIV, subsequent to this, the host cell undergoes programmed cell death, apoptosis of T cells. However, in other retroviruses, the host cell remains intact as the virus buds out of the cell.

Some eukaryotic cells contain an enzyme with reverse transcription activity called telomerase. Telomerase is a reverse transcriptase that lengthens the ends of linear chromosomes. Telomerase carries an RNA template from which it synthesizes DNA repeating sequence, or "junk" DNA. This repeated sequence of DNA is important because, every time a linear chromosome is duplicated, it is shortened in length. With "junk" DNA at the ends of chromosomes, the shortening eliminates some of the non essential, repeated sequence rather than the protein-encoding DNA sequence farther away from the chromosome end. Telomerase is often activated in cancer cells to enable cancer cells to duplicate their genomes indefinitely without losing important protein-coding DNA sequence. Activation of telomerase could be part of the process that allows cancer cells to become *immortal*.

Promoters

A promoter is a region of DNA that initiates transcription of a particular gene. Promoters are located near the transcription start sites of genes, on the same strand and upstream on the DNA (towards the 5' region of the antisense strand). Promoters can be about 100–1000 base pairs long.

For transcription to take place, the enzyme that synthesizes RNA, known as RNA polymerase, must attach to the DNA near a gene. Promoters contain specific DNA sequences such as response elements that provide a secure initial binding site for RNA polymerase and for

proteins called transcription factors that recruit RNA polymerase. These transcription factors have specific activator or repressor sequences of corresponding nucleotides that attach to specific promoters and regulate gene expression.

In bacteria

The promoter is recognized by RNA polymerase and an associated sigma factor, which in turn are often brought to the promoter DNA by an activator protein's binding to its own DNA binding site nearby.

In eukaryotes

The process is more complicated, and at least seven different factors are necessary for the binding of an RNA polymerase II to the promoter.

Promoters represent critical elements that can work in concert with other regulatory regions (enhancers, silencers, boundary elements/insulators) to direct the level of transcription of a given gene.

Promoter elements

Core promoter – the minimal portion of the promoter required to properly initiate transcription ^[3]

- Includes the transcription start site (TSS) and elements directly upstream
- A binding site for RNA polymerase
- RNA polymerase I: transcribes genes encoding ribosomal RNA
- RNA polymerase II: transcribes genes encoding messenger RNA and certain small nuclear RNAs and microRNA
- RNA polymerase III: transcribes genes encoding transfer RNAs and other small RNAs
- General transcription factor binding sites, e.g. TATA box
- Proximal promoter the proximal sequence upstream of the gene that tends to contain primary regulatory elements
- Approximately 250 base pairs upstream of the start site
- Specific transcription factor binding sites
- Distal promoter the distal sequence upstream of the gene that may contain additional regulatory elements, often with a weaker influence than the proximal promoter

• Anything further upstream (but not an enhancer or other regulatory region whose influence is positional/orientation independent)

• Specific transcription factor binding sites

Operator

• an operator is a segment of DNA to which a transcription factorbinds to regulate gene

expression. The transcription factor is a repressor, which can bind to the operator to prevent transcription.

• The main operator (O2) in the classically defined *lac* operon is located slightly downstream of the promoter. Two additional operators, O1 and O3 are located at -82 and +412, respectively.

Mechanism

- The repressor protein physically obstructs the RNA polymerase from transcribing the genes.
- An inducer (small molecule) can displace a repressor (protein) from the operator site (DNA), resulting in an uninhibited operon.
- Alternatively, a corepressor can bind to the repressor to allow its binding to the operator site. A good example of this type of regulation is seen for the trp operon.

Terminator

transcription terminator is a section of nucleic acid sequence that marks the end of a gene or operon in genomic DNA during transcription. This sequence mediates transcriptional termination by providing signals in the newly synthesized mRNA that trigger processes which release the mRNA from the transcriptional complex. These processes include the direct interaction of the mRNA secondary structure with the complex and/or the indirect activities of recruited termination factors. Release of the transcriptional complex frees RNA polymerase and related transcriptional machinery to begin transcription of new mRNAs.

Two classes of transcription terminators, Rho-dependent and Rho-independent, have been identified throughout prokaryotic genomes. These widely distributed sequences are responsible for triggering the end of transcription upon normal completion of gene or operon transcription, mediating early termination of transcripts as a means of regulation such as that observed in transcriptional attenuation, and to ensure the termination of runaway transcriptional complexes

that manage to escape earlier terminators by chance, which prevents unnecessary energy expenditure for the cell.

Rho-dependent terminators

Rho-dependent transcription terminators require a protein called Rho factor, which exhibits RNA helicase activity, to disrupt the mRNA-DNA-RNA polymerase transcriptional complex. Rho-dependent terminators are found in bacteria andphage. The Rho-dependent terminator occurs downstream of translational stop codons and consists of an unstructured, cytosine-rich sequence on the mRNA known as a Rho utilization site (*rut*) for which a consensus sequence has not been identified, and a downstream transcription stop point (*tsp*). The *rut* serves as a mRNA loading site and as an activator for Rho; activation enables Rho to efficiently hydrolyze ATP and translocate down the mRNA while it maintains contact with the rut site. Rho is able to catch up with the RNA polymerase, which is stalled at the downstream *tsp* sites. Contact between Rho and the RNA polymerase complex stimulates dissociation of the transcriptional complex through a mechanism involvingallosteric effects of Rho on RNA polymerase.

Rho-independent terminators

Intrinsic transcription terminators or Rho-independent terminators require the formation of a selfannealing hairpin structure on the elongating transcript, which results in the disruption of the mRNA-DNA-RNA polymerase ternary complex. The terminator sequence contains a 20 basepair GC-rich region of dyad symmetry followed by a short poly-T tract or "T stretch" which is transcribed to RNA to form the terminating hairpin and a 7-9 nucleotide "U tract" respectively. The mechanism of termination is hypothesized to occur through a combination of direct promotion of dissociation through allosteric effects of hairpin binding interactions with the RNA polymerase and "competitive kinetics". The hairpin formation causes RNA polymerase stalling and destabilization, leading to a greater likelihood that dissociation of the complex will occur at that location due to an increased time spent paused at that site and reduced stability of the complex. Additionally, the elongation protein factor NusA interacts with the RNA polymerase and the hairpin structure to stimulate transcriptional termination.

Termination in eukaryotes

In eukaryotic transcription of mRNAs, terminator signals are recognized by protein factors that are associated with the RNA polymerase II and which trigger the termination process. Once the poly-A signals are transcribed into the mRNA, the proteinscleavage and polyadenylation specificity factor (CPSF) and cleavage stimulation factor (CstF) transfer from the carboxyl terminal domain of RNA polymerase II to the poly-A signal. These two factors then recruit other proteins to the site to cleave the transcript, freeing the mRNA from the transcription complex, and add a string of about 200 A-repeats to the 3' end of the mRNA in a process known as polyadenylation. During these processing steps, the RNA polymerase continues to transcribe for several kilobases and eventually dissociates from the DNA and downstream transcript through an unclear mechanism; there are two basic models for this event known as the torpedo and allosteric models.

Torpedo model

After the mRNA is completed, the residual RNA strand remains in association with the DNA template and the RNA polymerase II, continuing to be transcribed. XRN2 (5'-3' Exoribonuclease 2), a RNase, attaches to the carboxyl terminal domain of RNA polymerase II and proceeds to degrade the uncapped residual RNA from 5' to 3' until it reaches the RNA pol II. 5' cap refers to a modified guanine added to the front of mRNA for protection from RNase. 3' poly(A) tail is added to the end of a mRNA strand for protection from exonucleases. Similar to Rho-dependent termination, XRN2 triggers dissociation of RNA polymerase II by either pushing the polymerase off of the DNA template or pulling the template out of the RNA polymerase. The entire mechanism remains unclear.

Allosteric model

RNA polymerase normally is capable of transcribing DNA into single-stranded mRNA efficiently. However, upon transcribing over the poly-A signals on the DNA template, a conformational shift is induced in the RNA polymerase from the proposed loss of associated proteins from its carboxyl terminal domain. This change of conformation reduces RNA polymerase'sprocessivity making the enzyme more prone to dissociating from its DNA-RNA substrate. In this case, termination is not completed by degradation of mRNA but instead is mediated by limiting the elongation efficiency of RNA polymerase and thus increasing the likelihood that the polymerase will dissociate and end its current cycle of transcription.

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Enhancer

An enhancer is a short region of DNA that can be bound with proteins (namely, the trans-acting factors, much like a set of transcription factors) to enhance transcription levels of genes (hence the name) in a gene cluster. While enhancers are usually cis-acting, an enhancer does not need to be particularly close to the genes it acts on, and need not be located on the same chromosome.

In eukaryotic cells the structure of the chromatin complex of DNA is folded in a way that functionally mimics the supercoiled state characteristic of prokaryotic DNA, so that although the enhancer DNA is far from the gene in regard to the number of nucleotides, it is geometrically close to the promoter and gene. This allows it to interact with the general transcription factors and RNA polymerase II. An enhancer may be located upstream or downstream of the gene that it regulates.

Furthermore, an enhancer does not need to be located near to the transcription initiation site to affect the transcription of a gene, as some have been found to bind several hundred thousand base pairs upstream or downstream of the start site. **Enhancers do not act on the promoter region itself, but are bound by activator proteins**. These activator proteins interact with the mediator complex, which recruits polymerase II and the general transcription factors which then begin transcribing the genes. Enhancers can also be found within introns. An enhancer's orientation may even be reversed without affecting its function. Additionally, an enhancer may be excised and inserted elsewhere in the chromosome, and still affect gene transcription. That is the reason that intron polymorphisms are checked though they are not translated.

Silencers

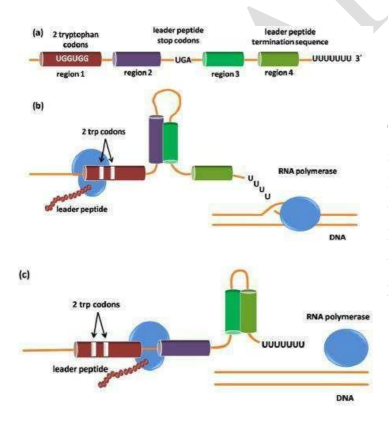
In genetics, a **silencer** is a DNA sequence capable of binding transcription regulation factors, called repressors. DNA contains genes and provides the template to produce messenger RNA (mRNA). That mRNA is then translated into proteins that activate or inactivate gene expression in cells. When a repressor protein binds to the silencer region of DNA, RNA polymerase is prevented from transcribing the DNA sequence into RNA. With transcription blocked, the translation of RNA into proteins is impossible. Thus, silencers prevent genes from being expressed as proteins.

RNA polymerase, a DNA-dependent enzyme, transcribes the DNA sequences, called nucleotides, in the 3' to 5' direction while the complementary RNA is synthesized in the 5' to 3' direction. RNA is similar to DNA, except that RNA contains uracil, instead of thymine, which forms a base pair with adenine. An important region for the activity of gene repression and expression found in RNA is the 3' untranslated region. This is a region on the 3' terminus of RNA that will not be translated to protein but includes many regulatory regions.

Not much is yet known about silencers but scientists continue to study in hopes to classify more types, locations in the genome, and diseases associated with silencers.

Attenuation

Premature termination of primary transcript in the leader region i.e. before the first structural genes is called attenuation. Attenuation is carried out by attenuator, a sequence within leader region of the tryptophan operon, (Fig 2). At this site, choice is made by RNA polymerase either to terminate or continue transcription. Mutants with small deletions in this region produce tryptophan synthesizing enzymes even in the presence of tryptophan.



Termination of Transcription regulated by attenuation (a) Stemloop structures of the trp operon in the mRNA; (b) Low level of trp full length mRNA made; (c) High level transcription of the trp operon is prematurely halted

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Expression of genes

• DNA serves two major functions:

1. Encoding structural information that can be converted into RNA and (usually), hence into protein sequence.

2. Encoding regulatory signals that allow certain proteins to decide where to begin or terminate reading DNA

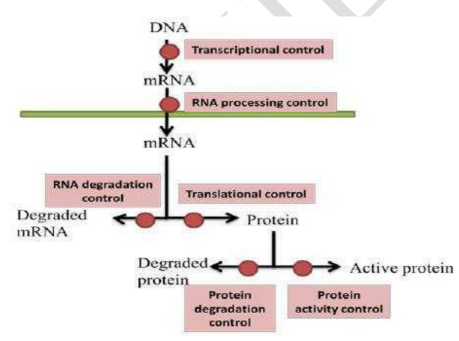
• The "*central dogma*" of molecular biology: information flows from nucleic acids to proteins, not the reverse:

DNA ----> RNA ----> polypeptides (proteins) Transcription = the process of making RNA

from DNA templates

• Translation = the process of making polypeptides

Control of gene expression:



Points of control of gene expression The eukaryotic gene expressions in a cell are controlled at about six stages

• Transcriptional control decides when and how frequently a given gene be transcribed. It enables the cell to check any discrepancy in the quantity of transcriptomes generated.

• RNA processing regulates how the RNA transcript is spliced or otherwise processed (applicable only to eukaryotes), which have split genes consisting of exons and introns.

- Translational control decides which mRNAs in the cytoplasm are to be translated by ribosomes.
- mRNA degradation control selectively destabilize some mRNA molecules in the cytoplasm.
- Protein activity control selectively activates, inactivates, degrades, or compartmentalizes

specific protein molecules after they have been synthesized. (applicable only to eukaryotes)

Regulatory elements:

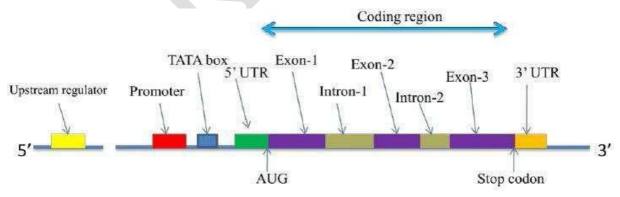
Gene expression is a complex multi-step process. Amongst various steps involved, transcription initiation is a vital key point in controlling gene expression.

Fundamental elements that regulate the process of transcription are:

- cis-acting elements comprising of special DNA sequences.
- trans-acting elements comprising of regulatory proteins.
- ° DNA -protein interaction.
- Protein-protein interaction
- RNA polymerase

cis-acting elements:

In Latin "cis", means "same side as". Cis acting elements are thus a region of DNA or RNA that regulates the expression of genes located on the same molecule. The cis-regulatory elements are often binding sites for one or more trans-acting factors. Cis-element may be located upstream to the coding sequence of the gene it controls (in the promoter region or further upstream 5'), in an intron, or 3' to the gene's coding sequence, either in the untranslated or untranscribed region.



Structure of eukaryotic gene with upstream regulatory elements

Examples of cis-acting elements:

Prokaryotic systems:

 \circ Promoter is a DNA sequence where RNA polymerase binds to initiate transcription. There are two promoter sequences in prokaryotic systems known as -10 (Pribnow box or Pribnow-Schaller box) and -35 sequences.

• Operator is regulatory sequence of DNA located immediately upstream of the structural gene that controls transcription of an operon.

• Inducers are located upstream of promoter region.

• Downstream regulatory sequences comprise of GC-rich inverted repeats followed by four adenine (A) residues signal the termination of transcription.

Trans-acting elements:

The protein factors which regulate the expression of gene by binding to cis acting DNA sequence are termed as trans-acting elements. Trans-acting molecules generally have two domains: DNA binding domain (which binds to cis elements) and protein binding domain (required for activation or suppression of transcription). Transcription initiation is a tightly regulated process controlled by trans-acting elements both in prokaryotes as well as eukaryotes.

DNA – Protein interactions: Gene regulatory proteins and the transcription factors are capable of binding to the DNA based on the interaction of amino acids of the protein with the nucleotides of the DNA. The regulation is implemented through various interactions between cis-acting elements and trans-acting factors. Examples of some DNA binding proteins are:

• Eukaryotic TATA-binding protein

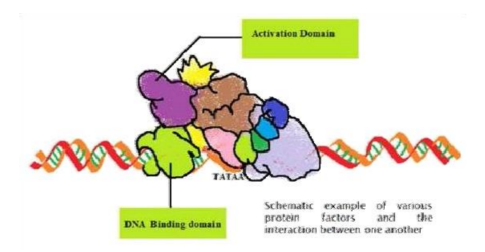
 $\circ \sigma$ subunit of bacterial RNA polymerase etc.

There are four types of structures of DNA binding proteins,

- 1. Zinc finger proteins
- 2. Helix loop Helix protein
- 3. Leucine zipper proteins
- 4. Homeodomain proteins

Protein – Protein interactions:

Protein-protein interaction is present in both prokaryotes and eukaryotes. The external signals affect the gene expression with the help of such interaction. Proteins interact with each other to form a homo or hetero-dimers before binding to DNA molecules. Eg. transcription initiation factors interacts with TATA box binding proteins (TBP) to activate transcription of a gene.



Interaction between various protein factors

RNA Polymerase:

• A single RNA polymerase is responsible for transcribing all types of RNA in prokaryotic system.

• However, eukaryotes have three different RNA polymerases, which have been found to specialize in the synthesis of various types of RNA:

• RNA polymerase I (Pol I) -transcribes rRNA (ribosomal RNA) genes.

• RNA polymerase II (Pol II) - transcribes protein-coding genes or mRNA (messenger RNA).

• RNA polymerase III (Pol III) - transcribes other functional RNA genes (e.g., tRNA).

• In eukaryotes, transcription occurs inside the nucleus. All the enzymes responsible for

translation are present in the cytosol therefore the transcripts formed then move out of the nucleus through nuclear pores into the cytosol (the liquid phase of the cytoplasm), where translation occurs. Organelle genomes (like the mitochondrial genome) are transcribed within the organelle (the mitochondria) and translation is also within the organelle (the mitochondria).

• Since prokaryotes have no nucleus, the step involving the movement of transcripts from nucleus to cytoplasm does not take place, and translation can take place immediately in the cytoplasm, directly on the growing transcript.

Possible questions

- 1. What are RNA polymerases?
- 2. Discuss on the role of promtors in transcription.
- 3. What is 5' capping?
- 4. Elaborate on the mechanism of transcription in eukaryotes.
- 5. Describe about the mechanism of transcription in prokaryotes
- 6. Explain the different types of RNA, its structure and role.
- 7. Explain the various elongation factors involved in transcription?
- 8. Write about the role of tRNA and mRNA in transcription.
- 9. Explain about the transcription process in prokaryotes.
- 10. Write in detail on RNA splicing.
- 11. Write about the role of transcription factors.
- 12. Give a detailed study on RNA processing in Transcription.

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UNIT IV

Regulation of gene expression and translation: Regulation of gene expression in prokaryotes: Operon concept (inducible and repressible system), Genetic code and its characteristics, Prokaryotic and eukaryotic translation: ribosome structure and assembly, Charging of tRNA, aminoacyl tRNA synthetases, Mechanism of initiation, elongation and termination of polypeptides, Fidelity of translation, Inhibitors of translation, Posttranslational modifications of proteins.

S.No	UNIT IV (Total No. of hours - 11 hours)	Reference Page No	
1	Regulation of gene expression – Overview	T2 410-415	
2	Regulation of gene expression in Prokaryotes	– R2 367-378	
3	Operon concept – Inducible and repressible		
4	Genetic Code and its Characteristics	T1 461-469	
5	Prokaryotic and Eukaryotic Translation	T1 483-527	
6	Ribosome structure assembly	R2 443-450	
7	Charging tRNA, aminoacyl tRNA synthetse		
8	Mechanism of Initiation, Elongation and Termination		
9	Fidelity of translation		
10	Post-translational Modifications		
11	Unit test and Discussions	-	

Reference

- 1. T1. Karp, G. (2013). Cell and Molecular Biology: Concepts and Experiments (7th ed). Hoboken, US: John Wiley & Sons. Inc.
- T2. Watson, J.D., Baker T.A., Bell, S.P., Gann, A., Levine, M., & Losick, R. (2008). Molecular Biology of the gene (6th edition). Cold spring Harbour Lab. Press, Pearson Pub.
- 3. R2. David Friefielder (2006). Molecular Biology (2006). Narosa Publishing house

GENE REGULATION

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Each cell expresses, or turns on, only a fraction of its genes. The rest of the genes are repressed, or turned off. The process of turning genes on and off is known as gene regulation. Gene regulation also allows cells to react quickly to changes in their environments. Gene regulation can occur at any point during gene expression, but most commonly occurs at the level of transcription (when the information in a gene's DNA is transferred to mRNA). Signals from the environment or from other cells activate proteins called transcription factors. These proteins bind to regulatory regions of a gene and increase or decrease the level of transcription. By controlling the level of transcription, this process can determine the amount of protein product that is made by a gene at any given time.

Introduction

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E. coli lives in colon. It has a metabolic pathway that allows for the synthesis of the amino acid tryptophan (Trp). o This pathway starts with a precurser molecule and proceeds through five enzyme catalyzed steps before reaching the final Trp product.

It is important that *E. coli* be able to control the rate of Trp synthesis because the amount of Trp available from the environment varies considerably.

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If we eat a meal with little or no Trp, the *E. coli* in our gut must compensate by making more.

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If we eat a meal rich in Trp, *E. coli* doesn't want to waste valuable resources or energy to produce the amino acid because it is readily available for use.

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Therefore, *E. coli* uses the amount of Trp present to regulate the pathway. o If levels are not adequate, the rate of Trp synthesis is increased.

If levels are adequate, the rate of Trp synthesis is inhibited.

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The first method works to decrease the synthesis of Trp by inhibiting the first enzyme in the pathway, preventing the rest of the pathway from proceeding.

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What inhibits the first enzyme? Trp does! \Box The more Trp in the cell, the more that can

bind to the first enzyme and prevent it from catalyzing the first step.

If the genes for the enzymes are not transcribed to mRNA, then translation to the

enzymes cannot occur.

Without enzymes, there is no Trp synthesis.

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This method of control is called regulation of gene expression because control is taking place at the genetic level. This method of control will now be examined in detail.

Trp Operon

The genes for the five enzymes in the Trp synthesis pathway are clustered on the same chromosome in what is called the Trp operon.

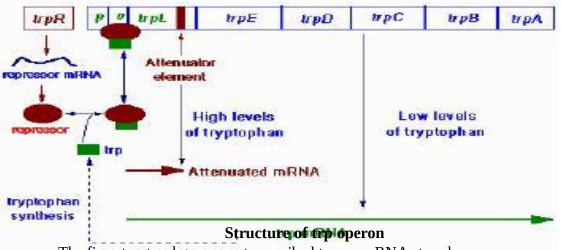
• The Trp operon has three components: o Five Structural Genes:

□ These genes contain the genetic code for the five enzymes in the Trp synthesis pathway

o One Promoter: DNA segment where RNA polymerase binds and starts transcription

o One Operator: DNA segment found between the promoter and structural genes. It determines if transcription will take place. If the operator is turned "on", transcription will occur.

• When nothing is bonded to the operator, the operon is "on". o RNA polymerase binds to the promoter and transcription is initiated.



o The five structural genes are transcribed to one mRNA strand.

o The mRNA will then be translated into the enzymes that control the Trp synthesis pathway.

• The operon is turned "off" by a specific protein called the repressor. o The repressor is a product of the regulator gene which is found some distance from the operon.

Transcription of

the regulator produces mRNA which is translated into the repressor.

The repressor is inactive in this form and cannot bind properly to the operator with this conformation.

o To become active and bind properly to the operator, a co-repressor must associate with the repressor. The corepressor if does is not strantist drop with the size of the siz

□ The more Trp available, the more that can associate with repressor molecules.

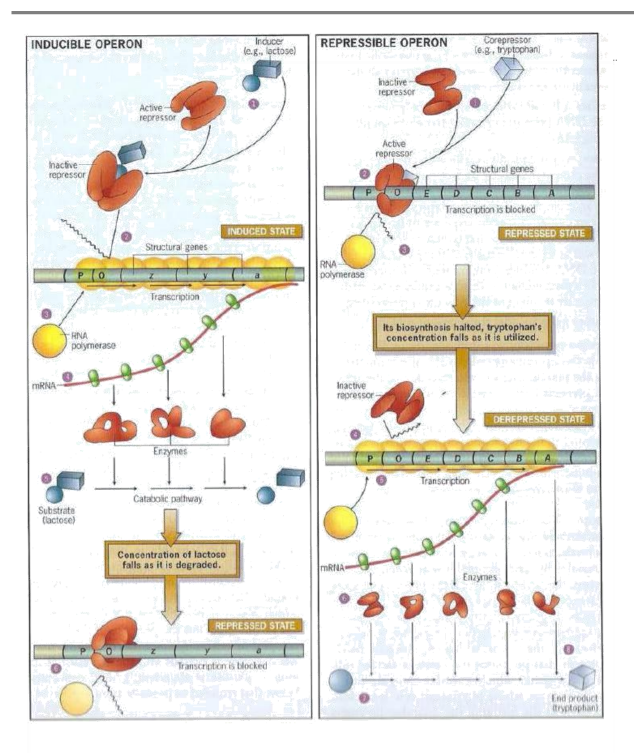
o An active repressor binds to the operator blocking the attachment of RNA polymerase to the promoter.

o Without RNA polymerase, transcription and translation of the structural genes can't occur and the enzymes needed for Trp synthesis are not made.

Repressible vs Inducible Systems

• The Trp pathway is anabolic as Trp is being synthesized. The Trp and other regulated anabolic pathways are usually repressible because the system can be repressed by an overabundance of the end product. o The end product, Trp, in this case, decreases or stops the transcription of the enzymes necessary for its production.

• Regulated catabolic pathways, on the other hand, are usually inducible because the pathway is stimulated rather than inhibited by a specific molecule. An example of an inducible system is lactose metabolism.



Inducible and repressible operons

The lac Operon

• The genes that code for the enzymes needed for lactose catabolism are clustered on the same chromosome in what is called the *lac* operon.

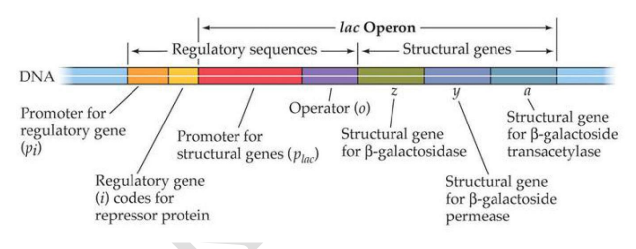
• The Lac operon has three components: o Three Structural Genes: \Box These contain the genetic code for the three enzymes in the lac catabolic pathway

o One Promoter:
DNA segment where RNA polymerase binds and starts transcription

o One Operator:
DNA segment found between the promoter and structural genes.

□ It determines if transcription will take place.

I If the operator in turned "on", transcription will occur.



The lac operon

• As in the Trp operon, the Lac operon is turned "off" by a specific protein called the repressor. o The repressor is the product of the regulator gene which is found outside the operon.

□ Transcription of the regulator produces mRNA which is translated into the repressor.

o But unlike the Trp operon, the repressor is active in this form and does not require a corepressor.

o The active repressor binds to the operator blocking the advancement of RNA polymerase to the structural genes.

o Without RNA polymerase, transcription and translation of the genes can't occur and the enzymes needed for Lac metabolism are not made.

• What turns the Lac operon "on"? Lactose does! o This makes sense because the cell only needs to make enzymes to catabolize lactose if lactose is present.

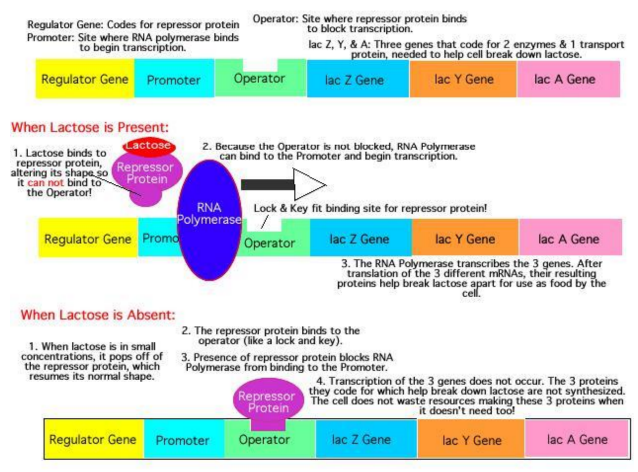
• Allolactose binds to the repressor and alters its conformation so that it can't bind to the operator.

• RNA polymerase can now start transcription. o The three structural genes are

transcribed to one mRNA strand.

o The mRNA will then be translated into the enzymes that control lactose catabolism.

• In this sense, allolactose is an inducer.



The *lac* operon: a model of gene regulation in prokaryotic

cells Negative vs Positive Control

• While the Trp operon is an example of repressible gene regulation and the Lac operon is an example of inducible gene regulation, both are examples of negative control of genes because

both operons are shut "off" by an active repressor.

• Gene regulation would be positive; on the other hand, if an activator molecule turned the operon "on".

• The Lac operon is also an example of a positive control system and is turned on by the cAMP-CAP complex, as described below:

• *E. coli* can be described as a fussy eater. o Its first choice at every meal is glucose because glucose supplies maximum energy for growth.

o Therefore, E. coli will only metabolize lactose if concentrations of glucose are low.

• For this to work, there must be a signal to tell the Lac operon that glucose is not available and to start transcribing the genes to metabolize lactose. o This signal is a small molecule called cyclic AMP (cAMP). □ The amount of cAMP present in a cell is inversely proportional to the amount of glucose present.

 \Box As a result, the absence of glucose results in an increase in cAMP in the cell.

• The following describes the situation where there is lactose but no glucose available to the cell: o No glucose means high levels of cAMP.

o cAMP binds to a molecule known as CAP.

o CAP, when in association with cAMP, can bind to the promoter at the CAP binding site.

o Here, the cAMP-CAP complex stimulates transcription by helping RNA polymerase bind to the promoter. \Box RNA polymerase has a weak affinity for the Lac promoter and will not bind without this help.

o Remember with lactose present so is allolactose. \Box Allolactose binds to the repressor and prevents it from binding to the operator.

 \Box Therefore, transcription and translation of the genes can occur.

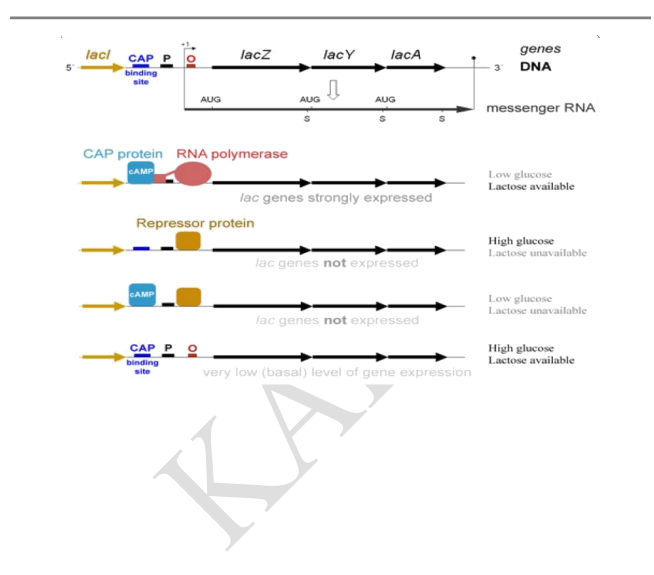
o The following depicts what happens when glucose and lactose are both present for *E*. *coli* to metabolize: \Box With glucose present, there is very little or no cAMP.

 \Box It cannot bind to the CAP binding site.

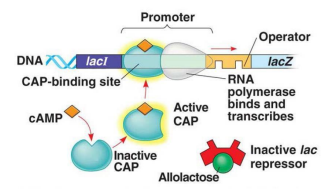
□ Without this complex, RNA polymerase cannot bind to the promoter and transcription cannot occur.

 \Box Even though allolactose is present and blocks the action of the repressor, there is no

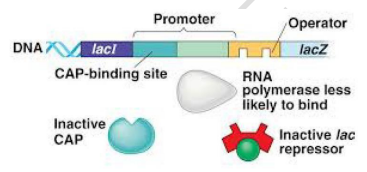
transcription of the lac genes because glucose is present.



Overall structural elements of Lac Operon



Lactose present, glucose scarce (cAMP level high): abundant lac mRNA synthesized



Lactose present, glucose present (cAMO level low): little lac mRNA synthesized

The Genetic code

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Since there are 20 different kinds of amino acids in proteins and only four kinds of nucleotides in DNA, the relationship between the gene and its most elementary functional product, i.e., between DNA and protein, can hardly be interpreted through a code of one nucleotide = one amino acid.

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A coding sequence of two nucleotides for one amino acid, or a doublet code, would produce only 16 possible coding combinations, or codons. Codons are group of nucleotides that specifies one amino acid. By the **genetic code** (George Gamov, 1954), we mean, a collection of base sequences (codons) that correspond to each amino acid and to translation signals.

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A codon size of three nucleotides for one amino acid are triplet codon seems more likely, since it produces 64 possible codons, however, only 20 amino acids need to be coded, 44 codons in a triplet code seem to be superfluous. To account for the excess of codons beyond the necessary 20, we can suppose that more than one codon can code for a particular amino

acid. For example, if each kind of amino acid were coded by three different possible codons, 60 possible codons would be accounted for.

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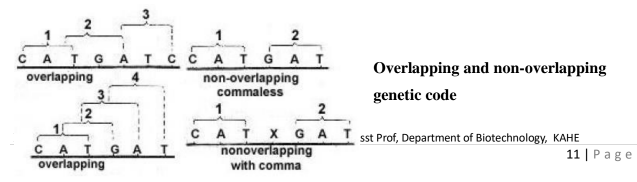
A code in which there is more than one codon for the same amino acid, is called degenerate. It is also possible that some or all of the codons in excess of 20 do not code for any amino acid and are therefore nonsense codons.

	Second letter								
		U	С	А	G				
etter	U	UUU UUC UUA UUG Leu	UCU UCC UCA UCG	UAU UAC UAA Stop UAG Stop	UGU UGC UGA Stop UGG Trp	U C A G			
	с	CUU CUC CUA CUG	CCU CCC CCA CCG	CAU CAC CAA CAA CAG GIn	CGU CGC CGA CGG	U C A G	letter		
First letter	A	AUU AUC AUA AUG Met	ACU ACC ACA ACG	AAU AAC AAA AAG Lys	AGU }Ser AGC }Arg AGA }Arg	UCAG	Third letter		
	G	GUU GUC GUA GUG	GCU GCC GCA GCG	GAU GAC GAA GAG Glu	GGU GGC GGA GGG	DCAG	1		

The genetic code has following general properties, mostly applicable to the genes of all the organisms:

Genetic code is triplet

As discussed earlier, singlet and doublet codons cannot form 20 combinations, which is the minimum requirement; therefore triplet codon is a necessity, so that all the amino acids must be coded. Genetic code is non-overlapping. During translation, the codons are read one after another, in a sequence. One base of a codon is not used by the other codons. Therefore, if there are six bases, they will code for two amino acids only. e.g.; in case of non-overlapping, a gene sequence of UUUCCC only two amino acids will be coded, phenylalanine (UUU) and proline (CCC), whereas for an overlapping code, more than two amino acids could be coded, phenylalanine (UUU), serine (UCC) and proline (CCC).



Genetic code is commaless

The bases are read one after the other in the codons, i.e., no bases or codons are reserved for punctuation or comma. When the first amino acid is coded, the second will be coded by the next three bases immediately, and no base will be wasted to serve as a comma. Once the translation begins, the codons are read one after the other with no break or demarcating signals in between them.

Genetic code is non-ambiguous

Each codon has a particular amino acid for coding, and it will code for that amino acid only. There is one to one relationship between codon to amino acid. However, there is an exception, AUG codes for methionine and GUG codes for valine, but if AUG is absent, then GUG codes for methionine, as starting codon for protein synthesis. In an ambiguous code, one codon can code for more than one amino acid. In certain rare cases, the genetic code is found to be ambiguous i.e, some codons codes for different amino acids under different

conditions, e.g. in streptomycin sensitive strain of E.coli the codon UUU normally codes for phenylalanine, but it may also code for isoleucine, serine when treated with streptomycin. This ambiguity is enhanced at high Mg ion concentration, low temperature and the presence of ethyl alcohol.

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Genetic code is degenerate. Since there are more codons than the amino acids, more than one codon may specify the same amino acid. Such different codons that specify the same amino acid are called as synonymous codons. e.g., UUU = UUC = phenylalanine.

 \succ

Genetic code has start/stop signals. Some codons are specially meant for initiation and termination of protein synthesis. e.g.: AUG codes for methionine, serves as initiation codon in eukaryotes and GUG in case of prokaryotes. Three codons UAG (**amber**), UAA (**ochre**), UGA (**opal**) are called as termination codons, because they terminate protein synthesis. Earlier, they were called as nonsense codons, because they do not code for any amino acid, but, since they are involved in termination of protein synthesis, they are called as termination codons are known as **signals** and this phenomenon is known as **punctuation**.

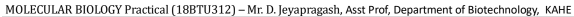
 $[\]triangleright$

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 \geq

Genetic code is polar. It means that the genetic code has a fixed start and termination ends, and is always read in a fixed direction, i.e. in $5' \rightarrow 3'$ direction and the polypeptide chain is synthesized in N \rightarrow C direction i.e., from amino group (NH2) to carboxylic group (COOH).

Genetic code is universal. The same genetic code is applicable to all organisms, from bacteria to man, i.e.; the codons have the same meaning in all the organisms. e.g., UUU = phenylalanine in bacteria, mouse, man and tobacco. The universality of the genetic code, however, does not mean that DNA base ratios must be similar in different species for genes specifying similar proteins. The fact that the code is degenerate enables many bases to be changed by mutation in a sequence of mRNA, but this mRNA could still produce the same amino acid sequence. In 1979 investigators began started DNA sequencing of mitochondrial DNA in humans, cattle and mice. During their experiments, they were surprised to learn that the genetic code used by the mitochondrial DNA was not the same as the universal genetic code. e.g. UGA, which is a non-sense codon, but it codes for tryptophan in mtDNA, AGG which codes for arginine is a non-sense codon in mtDNA. So, extra chromosomal DNA such as mtDNA and ctDNA do not come under the universality of the genetic code.



Wobble Hypothesis. Out of the 64 codons, three are involved in termination process. So, there are only 61 codons specifying the amino acids, and the cell should have 61 different types of tRNAs, each having a different anticodon for the recognition of codons. However, the actual number of tRNA is found to be much less than 61. This means that the anticodons of tRNA read more than one codon on the mRNA. **Crick** (1966) proposed a hypothesis to explain the degeneracy of the genetic code; the hypothesis is known as Wobble hypothesis.

According to this hypothesis, the major degeneracy occurs at the third position, i.e., the third codon is not important in base pairing, and the actual pairing occurs only in the first two codon-anticodon pairs. The base at the 5' end of the anticodon and the base at the 3' end of the codon form hydrogen bonds without any specificity. The third base is called as the **wobble base**. This wobble base of codon lacks specificity and the base in the first position of the anticodon is usually abnormal e.g., inosine, tyrosine, etc. These abnormal bases are able to pair up with more than one nitrogen base at the same position e.g., inosine (I) can pair up with A, C and U. The pairing between unusual base of tRNA and wobble base of mRNA is called **wobble pairing**.

Translation: the Synthesis of Proteins

Role of mRNA

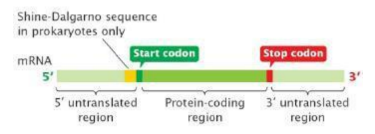
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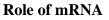
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• carries **codons** (3-nucleotide sequences) arranged in linear fashion that code for amino acids

• also carries signals needed to tell how to recognize ribosomes, start and stop signals for decoding protein

• **leader sequence** on small ribosome subunit binds to complementary sequence on mRNA, allows initial formation of RNA-ribosome complex.



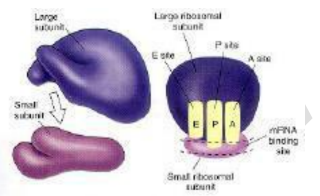


Role of ribosome

- two parts: a **small subunit** and a **large subunit**. These are separated except when attached to m-RNA
- ribosomes contain a set of ribosomal proteins and several types of ribosomal RNA (r-

RNA)

• ribosomes catalyze the formation of peptide bonds; intially thought to be due to protein activity (enzyme), but now known to be due to RNA catalytic activity (**ribozyme**).



Ribosome Subunits

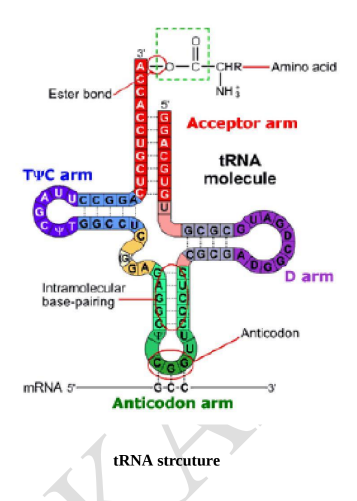
The smaller subunit its into a depression on the surface of the larger and. The A, F, and E sites on the ribosome play key roles in protein synthesis.

Ribosome

Role of tRNA

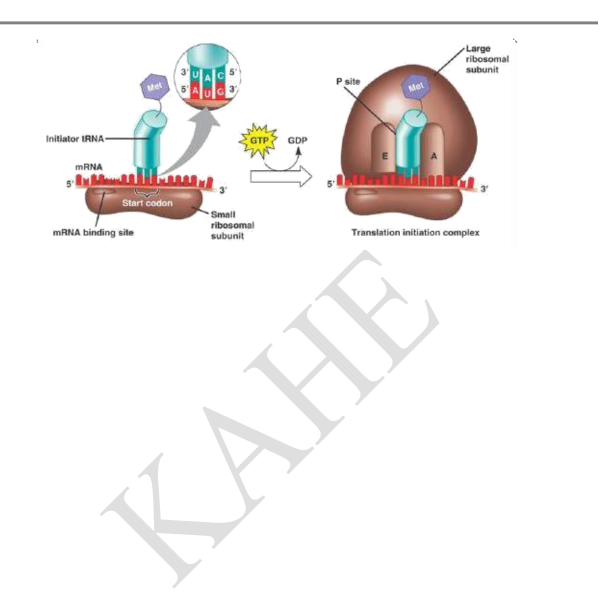
- structure: 4 loops, anticodon, AA binding site
- ~ 60 types in bacteria (>100 in mammals)
- only 73-93 nucleotides long
- some bases modified after transcription; like pseudouridine.
- extensive hairpin loops
- anticodon site: recognizes codon on mRNA
 - Activation of tRNA: adding amino acids o requires special enzyme: AA-tRNA activating enzymes
 - o ATP required, forms AA-AMP + PP, then AA-tRNA + AMP

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Initiation of Translation

- small ribosomal subunit initiates binding to mRNA
- locates 5' end of mRNA
- small subunit ribosome finds first AUG codon = start codon
- large ribosome binds
- tRNA carries the amino acid methionine to first position

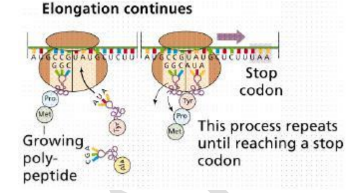


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Initiation of translation

Elongation of Translation

- 2 adjacent sites on ribosome: P (Peptide) and A (Amino Acid) site
- A site accepts a new tRNA-AA
- P site holds existing chain
- peptide transferred from P site tRNA to A-site AA
- enzyme activity is in ribosomal RNA (ribozyme)
- also required: Energy (GTP) and elongation factors



Elongation of translation

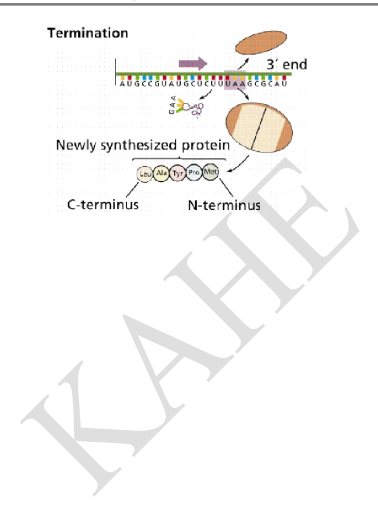
Termination of Translation

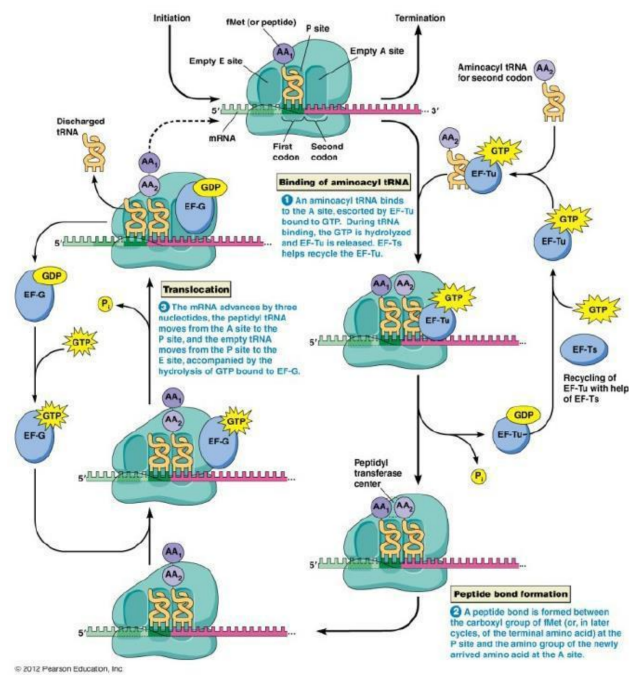
- reach a "stop codon" UAG, UAA, or UGA
- no t-RNAs bind
- instead, specific release factors required
- Net cost of translation: 4 phosphate bonds/amino acid added.

Termination of translation

NOTE: Typical RNA polymerization rate (~ 40 nts/second at 37 °C in bacteria - close to

translation rate of about ~15 aa/second).





The complete process of translation

Possible Questions

- 1. Write about post-translational modification of proteins
- 2. Describe in detail about the tryptophan operon
- 3. Explain the detail process of regulation of gene expression in prokaryotes.
- 4. Explain in detail about the process for protein synthesis.
- 5. Write about the ribosome structure and assembly.
- 6. Elaborate on various steps involved in DNA replication in prokaryotes.
- 7. Explain: (i) the features of genetic code, (ii). Poly-adenylation.
- 8. Explain in detail about *Lac* operon
- 9. Explain about Operon concept with an example.
- 10. Elaborate on fidelity of translation.

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UNIT V

DNA damage, repair and homologous recombination: DNA damage and repair: causes and types of DNA damage, mechanism of DNA repair: Photoreactivation, base excision repair, nucleotide excision repair, mismatch repair, trans-lesion synthesis, recombinational repair, nonhomologous end joining. Homologous recombination: models and mechanism.

S.No	UNIT V (Total No. of hours - 9 hours)	Reference page No
	DNA Damage – Overview	T1 591-596
	Causes and types of DNA Damage	R2 280-290
	Mechanism of DNA repair	R2 280-290
	Photoreactivation and their genes	R2 280-290
	Base Excision, nucleotide excision and repair mechanism	R2 280-290
	Mismatch repair and translational synthesis	R2 280-290
	Recombinational repair	R2 280-290
	Homologous end joining, Recombination models and Mechanisms	R2 280-290
	Unit test and discussion	-

Reference

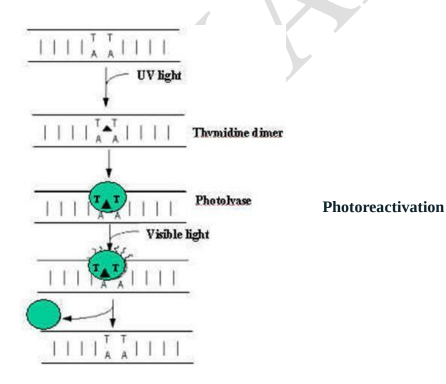
- R2. David Friefielder (2006). Molecular Biology (2006). Narosa Publishing house.
- T1. Karp, G. (2013). Cell and Molecular Biology: Concepts and Experiments (7th ed). Hoboken, US: John Wiley & Sons. Inc.

DNA REPAIR

It can be defined as **"any one of the cellular process that attempts to correct errors in cellular DNA which occurred during cell division or by external environment".** There are many cellular proteins and enzymes which help in repairing most of the errors by means of different mechanisms.

Photoreactivation

- Catalyzed by DNA-Photolyase (DPL)
- Reverses cyclobutyl pyrimidine dimers resulting from UV irradiation.
- It has low abundance (10 to 20 molecules per cell) in E. coli.
- DPL binds irradiated DNA 100x better than non-irradiated DNA.
- DPL contains a non-covalently associated chromophore (FADH or FADH2).
- The photochemical mechanism of photorepair has been proposed to be light-dependent redox reaction between the singlet excited state of FADH2 and the pyrimidine dimer.
- Photolyase has not yet been identified in placental mammals.



Aberrant Methylation

- Catalyzed by 6-O-methylguanine methyltransferase (6-O-MGM)
- 6-O-MGM recognizes 6-O-methylguanine in DNA and removes the methyl group, transferring the group to an amino acid on itself in a "suicide" mechanism.
- 6-O-MGM is encoded by the ada gene in E. coli and the MGMT gene in eukaryotes.
- Mice knocked out for MGMT are cancer-prone and sensitive to methylation agents

Common Features of Excision-Type Repair Pathways

- Recognition: Altered DNA is recognized and bound by a specific damage-recognition protein. This first step recruits other components required for the repair reaction.
- Excision: Damaged base(s), and in some cases adjacent nucleotides, are excised from the strand by exonucleases, resulting in a gapped DNA.
- Resynthesis: The gap is refilled by a DNA polymerase using the complementary strand as a template.

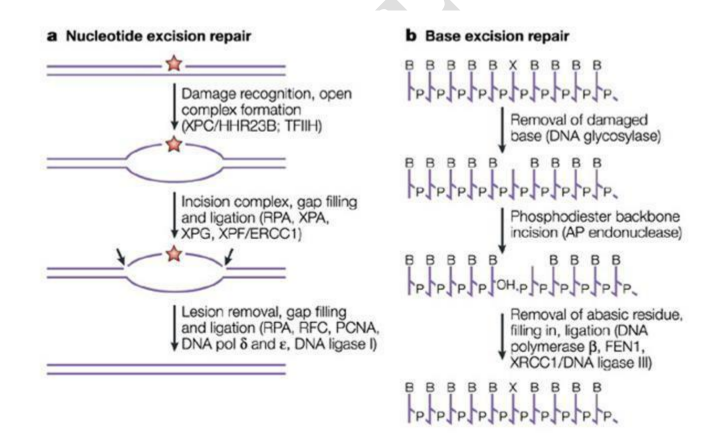
Base-Excision Repair (BER)

- Damaged bases are removed as free bases.
- BER primarily handles oxidative and alkylative damage.
- BER is thought to have an important role in aging.
- BER recognizes base deamination, oxidative damage, and other minor base modifications.
- Five gene products are required for BER: glycosylase, AP endonuclease, phosphodiesterase, DNA polymerase, and DNA ligase.
- DNA glycosylase recognizes the damaged base and removes it, generating an AP (apurinic, apyramidinic) site.
- AP endonuclease cleaves the phosphodiester bond, generating a single-strand break with a 5'terminal deoxyribophosphate moiety.
- The 5'-deoxyribophosphate is excised by action of a DNA phosphodiesterase.
- The resulting single-nucleotide gap is repaired by DNA polymerase β (beta).
- The resulting nick is sealed by DNA ligase.

• BER is relatively inefficient, due to the large number of peptides needed to recognize each damage type.

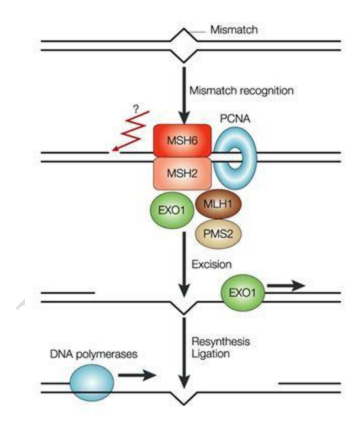
Nucleotide-Excision Repair (NER)

- Damaged bases are removed as oligonucleotides.
- NER is primarily responsible for removal of UV-induced damage and bulky adducts, but also removes ~20% of oxidative damage.
- Deficiencies cause many human disorders.
- Xeroderma Pigmentosum (both classical and variant) and Cockayne's syndrome are caused by defects in NER, resulting in various detrimental effects upon exposure to UV light.



DNA Mismatch Repair (MMR)

- Mismatches are removed as long oligonucleotides.
- MMR is primarily responsible for removal of replicative errors.
- Also prevents recombination of non-homologous sequences
- Deficient in many human cancers.
- MMR recognizes base-base mismatch and small insertion/deletion loops.



DNA mismatch repair

Error prone translation DNAsynthesis

When this pathway is active, DNA repair becomes significantly less accurate and a high mutation rate occurs. It is a part of cellular stress response to extensive DNA damage, also known as SOS response. The cell, at all cases had to survive, so DNA gets repaired, although it contains a lot of errors.

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Types of Mutations

NOTE: For all others examples cited below, the below given DNA sequence shall be considered as wild type:

- ATG CCG TGT CAG ATG TTC ------ DNA
- AUG CCG UGU CAG AUG UUC ----- mRNA

Met Pro Cys Gln met phe ----- Amino acid sequence

1. Synonymous / Silent Mutations

- 1. No alteration in polypeptide product of the gene
- 2. Single base pair substitution
- 3. Occur in the third position of a codon
- 4. Codes for the same amino acid
- 5. No alteration of the protein

Example: *Samesense Mutation*: a codon is changed to a different codon that specifies the same amino acid.

ATG CCG TGC CAG ATG TTC-----MUTATED DNA (compare with wild type)AUG CCG UGC CAG AUG UUC-----mRNA

Met Pro Cys Gln Met Phe ------ amino acid sequence

2. Non-Synonymous Mutations

- 1. Occur less frequently than synonymous mutations
- 2. Leads to alteration in the encoded polypeptide
- 3. Result in abnormal function® disease

Occur in one of three main ways:

- 1. Missense
- 2. Nonsense
- 3. Frameshift

Missence mutation: a codon is changed to a different codon that specifies a different amino acid.

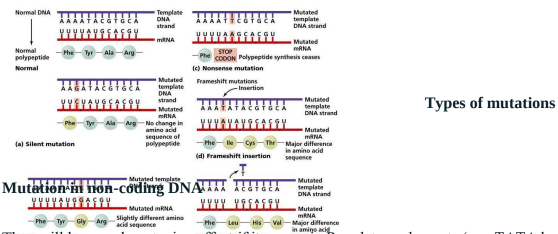
ATG CCG TGG CAG ATG TTC-----MUTATED DNAAUG CCG UGG CAG AUG UUC----- mRNAMetPro Trp Gln Met Phe ------ amino acid sequence

Nonsense Mutation: A codon that specifies an amino acid is changed to a stop codon. (termination codon), this mutation usually destroys the function of the gene product.

ATG CCG TGA CAG ATG TTC----- MUTATED DNAAUG CCG UGA CAG AUG UUC----- mRNAMet Pro STOP------ amino acid sequence

Frameshift Mutations:

One or 2 nucleotide pairs are inserted into or deleted from the molecule, causing an alteration of the *reading frame*. As the result of this shift, codons downstream of the insertion or deletion site specify an entirely new sequence of amino acids. Depending on where the insertion or deletion occurs in the gene, different effects can be generated. In addition to producing an entirely new polypeptide sequence immediately after the change, frameshift mutations usually produce a stop or termination codon within a short distance of the mutation. This codon terminates the already altered polypeptide chain. A frame shift in a gene specifying an enzyme usually result in a loss of enzyme activity. If the enzyme is an essential one, the effect on the organism can be disastrous.



The y-will-have a phenotyping effect if it occurs in Regulatory elements (e.g. TATA box) \rightarrow MOLECULAR BIOLOGY (18BTU302) – Mr. D. Jeyapragash, Asst Prof, Department of Biotechnology, KAHE

affect level of gene expression

1. Splicing of introns: highly conserved GT & AG at the end of introns

Either:

- a. coding sequences being lost
- b. intronic sequences being added to the mRNA

Functional effects of Mutation on the Protein

- 1. Loss of function
- 2. Gain of Function

Loss of Function

• Reduce in activity

In heterzygous state \rightarrow half normal levels of the protein product

• Or complete loss of the gene product

Gain of Function:

- Increased levels of gene expression
- Development of a new function(s) of the gene product

Chromosomal Mutations:

Chromosomal mutations are grouped into four broad types:

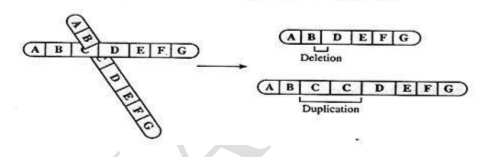
1. **Deletion**: are chromosomal changes in which one or more genes or segments of chromosomal DNA are lost.

A. Deletion mutation - loss of a chromosomal segment



2. Duplication: are chromosomal changes in which one copy or more copies of a gene are present on the same chromosome

> B. Deletion and Duplication mutations – exchange of chromosomal segments between homologous chromosomes that break at different points



3. Inversion: in which a segment of DNA is released and rotated 180 degrees before being reinserted into the DNA. If the inverted DNA segment carries part of the protein-coding sequence, the resulting protein would be drastically altered and most likely nonfunctional.

C. Inversion mutation – removal of a segment of chromosomal DNA and its reinsertion in the reverse orientation



4. Translocation: occur when a segment of DNA moves from one chromosome and inserted into a different nonhomologous chromosome. Translocation can also be reciprocal, that is two nonhomologous chromosome may break and trade pieces of DNA. Translocation mutations frequently cause

problems in meiosis and sometimes lead to aneuploidy (the gain or loss of chromosome)

DNA recombination

DNA recombination involves the exchange of genetic material either between multiple chromosomes or between different regions of the same chromosome. This process is generally mediated by homology; that is, homologous regions of chromosomes line up in preparation for exchange, and some degree of sequence identity is required.

- Homologous recombination occurs between two homologous chromosomes.
- Nonhomologous, or illegitimate recombination occurs between two different chromosomes, though the segment at recombination sites may be related.
- Site-specific recombination can result in integration of viral, bacterial, or plasmid DNA into a chromosome at a specific location (such as att).
- Replicative recombination results in sequence transposition, and is mediated by transposase enzymes.

Types of Homologous Recombination

- Reciprocal recombination results from two chromosomes exchanging the same amount of DNA.
- Non-reciprocal recombination results from two chromosomes exchanging different amounts of DNA.
- Intramolecular recombination results from recombination in chromosomal loops.
- Direct repeats cause plasmid formation from the intervening loop.
- \circ Inverted repeats cause the intervening loop to change direction.

• Double crossover results in exchanging a short segment of DNA, rather than a large segment of chromosome from the crossover point to the terminus.

Holliday Model for Homologous Recombination

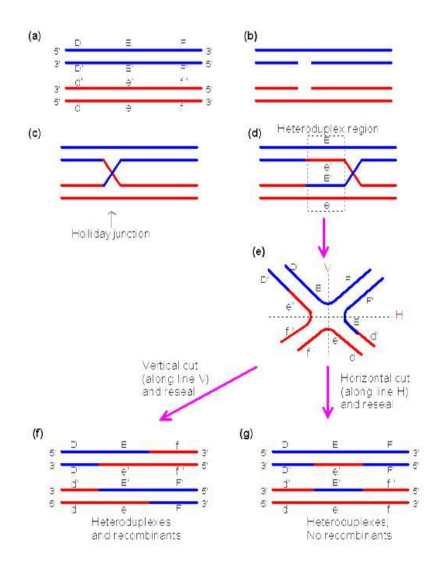
• Two DNA molecules with nicks induced by an endonuclease in strands of the same polarity can invade each other with free single strands.

• A Holliday junction, or Chi structure, is formed as a recombination intermediate, and is solidified by ligase sealing the nicks, uniting the two homologues.

• Branch migration allows exchange of material, since the loose area around the junction allows unzipping/rezipping.

• New nicks allow separation of recombined DNA and resolution of the recombination intermediate.

• The Holliday model assumes reciprocal and equal exchange of genetic material between DNA molecules.



Holliday model

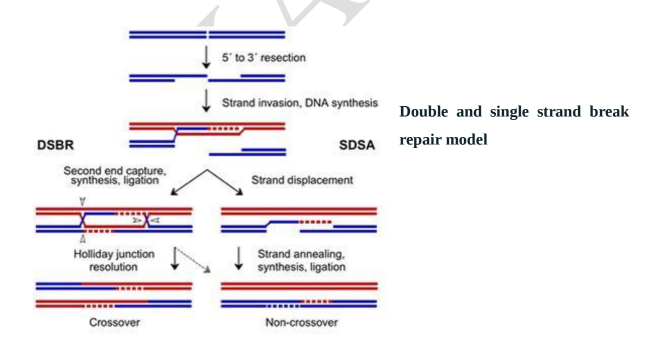
Double-Strand Break Repair Model (Szostak Model) for Recombination

- Double-strand breaks are introduced into one of the homologues.
- Broken ends are recessed by specific exonucleases which produce longer 3' single strands.
- The single strands can invade a region of homologous DNA.

• The invading end serves as a primer for a polymerase which extends it, unwidnding the template DNA and generating a D-loop. A Holliday junction is formed, and DNA synthesis proceeds until it reaches the opposing end of the recombining chromosome.

• The displaced DNA of the opposing end of the invading chromosome anneals to the template (invaded chromosome), forming a second Holliday junction by ligation of the strands. Holliday junctions can migrate prior to resolution with resolvase endonucleases.

- Resolution can cut the invading strands, resulting in a non-crossover event, since each chromosome has only a tiny fragment of DNA from the other chromosome.
- Crossover results from cutting the strands not participating in the recombination, generating a crossover by leaving a large segment of each chromosome attached to the other.
- The Szostak model accounts for non-reciprocal recombination.



MOLECULAR BIOLOGY (17BTU302) - Dr. J. Anburaj Factors Involved in Each Recombination Step

Page 12

• The Szostak model is an error-free pathway (since it can only occur in homologues), and is of major importance in yeasts and mammals.

- A 5' to 3' recession reaction is mediated by endonucleases.
- Strand invasion requires multiple Rad proteins, including DNA helicases and DNA endbinding proteins.
- New DNA synthesis requires DNA Pol δ and ϵ (delta and epsilon).

Repair of DSBs by Non-Homologous End Joining (NHEJ)

- NHEJ is an error-prone pathway, and is the major pathway in mammals for repair of DSBs.
- DNA-PKcs (DNA protein kinase, catalytic subunit) binds ends of the DSB (double-strand break).
- Endonucleases create blunt (no overhang) ends.
- Synapsis is achieved through microhomologies.

• Ends are ligated, and, due to end-blunting and other factors, small insertions and deletions are introduced. The errors are often of little consequence, since most DNA is non-coding.

Site specific recombination

Site-specific recombination, also known as conservative site-specific recombination, is a type of genetic recombination in which DNA strand exchange takes place between segments possessing only a limited degree of sequence homology. Site-specific recombinases

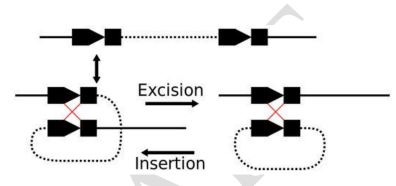
perform rearrangements of DNA segments by recognizing and binding to short DNA sequences (sites), at which they cleave the DNA backbone, exchange the two DNA helices involved and

rejoin the DNA strands. While in some site-specific recombination systems just a recombinase enzyme and the recombination sites is enough to perform all these reactions, in other systems a number of accessory proteins and/or accessory sites are also needed.

Site-specific recombination systems are highly specific, fast and efficient, even when faced with

complex eukaryotic genomes. They are employed in a variety of cellular processes, including bacterial genome replication, differentiation and pathogenesis, and movement of mobile genetic elements. For the same reasons, they present a potential basis for the development of genetic engineering tools.

Recombination sites are typically between 30 and 200 nucleotides in length and consist of two motifs with a partial inverted-repeat symmetry, to which the recombinase binds, and which flank a central crossover sequence at which the recombination takes place. The pairs of sites between which the recombination occurs are usually identical, but there are exceptions (e.g. attP and attB of λ integrase,)



Insertion and excision mediated by aligned Lox sites and the cre recombinase. Red X designates recombination.

Possible Questions

- 1. Explain about homologous recombination.
- 2. Write a short note on proteins involved in base excision repair.
- 3. Briefly explain about the different types of DNA damages.
- 4. Define Mutation. Explain in detail about different types of Mutations.
- 5. Write about the Nucleotide excision repair.
- 6. Describe about non homologues recombination.
- 7. Explain the detail process of DNA repair mechanism.
- 8. Write about the formation of Holliday model.
- 9. Write about the Mismatch repair occurring during DNA damage.
- 10. Explain in detail on the formation of thymine thymine and its repair mechanism.

The backbone of	Hydrogen bond	Phosphodiester	Ionic bond	Hydrophobic	Hydroge
nucleic acid structure is		bond		bond	n bond
The functional unit of	Genome	Gene	Nucleotide	Chromosome	Nucleotid
DNA is					e
Base pairing of DNA is	Arther kornberg	Chargaff's	Meselson	Mc clintock	Charhoff
based on rules			&Stahl		
In Hershey and Chase	Protein	RNA	DNA	Gene	DNA
experiment P32 is					
incorporated in					
formation of					
RNA contain uracil in	Adenine	Guanine	Cytosine	Thymine	Adenine
place of					
The proteins associated	Trypsin	Histones	Collagen	Elastin	Histones
with eukaryotic DNA			C		
are					
DNA double helix does	Antiparallel	Complementary	Major and	Uracil	Uracil
not have	configuration	base pairing	minor grooves		
The total genetic	Gene	Genome	Ribosome	Chromosome	Genome
information contained					
in a DNA is referred to					
as					
Helicases enzyme has	Rewinding of DNA	Polymerize of	Ligase of DNA	Unwinding of	Unwindi
function of		DNA	Liguse of Divit	DNA	ng of
		DIMI		DIMI	DNA
Histones are rich in	Arg & Lys	Lys & Gly	Arg & Glu	Arg & Gly	Arg &
	118 00 200				Lys
Fundamental unit of	Replisome	Nucleosome	Primosome	Chromosome	Nucleoso
DNA organization is	reprisonie	i (dereosonie	i innobonne		me
B-DNA form was	Watson and Crick	Hoogsteen	Chargaft	Wang and Rick	Watson
discovered by		110080000	Chargare		and Crick
A nucleotide in DNA is	Deoxyribose sugar.	Deoxyribose sugar	Deoxyribose	Deoxyribose	Deoxyrib
composed of	phosphate & nitrogen	& nitrogen base	sugar &	sugar	ose
•••mp======	base		phosphate	Sugar	sugar,
	ouse		phosphate		phosphat
					e &
					nitrogen
					base
In a nucleotide, the	Phosphodiester bond	Glycosidic bond	Hydrogen bond	Hydrophobic	Glycosidi
nitrogen base is joined	i nosphoulester bolid		riyurogen bollu	bond	c bond
to the sugar molecule				DOILG	c bolla
J. J					
by Change in gualactide	nonligation	recombination	namain	mutation	mutation
Change in nucleotide	replication	recombination	repair	mutation	mutation
sequence of a short					
region of genome is					
called	1	1 . 1	1	1.0	1
Eukaryotic DNA is	nucleotides	nucleosides	nucleosomes	nucleofomes	nucleoso
bound to histones to					mes
form					

In which direction does	3'-5'	5'-3'	5'	3'	5'-3'
DNA replication take	5.5	5 5	5	5	5 5
place?					
The ends of the linear	helicase	primase	gyrase	telomerase	telomeras
chromosomes are		F	8,		e
maintained by					
The experiments by	radioactive phage were	radioactive cells	radioactive	radioactive	radioacti
Hershey and Chase	found in the pellet	were found in the	sulfur was	phosphorus was	ve
helped confirm that	iouno in uno poneo	supernatant		found in the cell	phosphor
DNA was the hereditary		supurnation	cell		us was
material on the basis of					found in
					the cell
Which histone protein	H1	H2a	H2b	H4	H1
is not part of a core			-		
particle of nucleosome?					
r					
Heterochromatin is	Naked DNA form	extended	solenoid form	looped solenoid	Naked
DNA in which form?		nucleosome form		form	DNA
					form
Double strrand nature	Watson & crick	Arther kornberg	Stahl &	Mc clintock	Watson
of DNA proposed by		C	Meselson		& crick
An enzyme that	DNA polymerase	DNA ligase	RNA	Topoisomerase	DNA
catalyses		C	polymerase	1	ligase
phosphodiester linkages					C
in DNA strands is					
called					
In DNA, the genetic	purine bases	pymidine bases	purine &	Sugar	purine &
information resides		1.	pyrimidine	C	, pyrimidin
in bases			bases		e bases
Z-form of DNA is	Left handed	Right handed	Both right and	Non directional	Left
		C .	left handed		handed
DNA synthesis takes	G1 Phase	S-Phase	G 2 Phase	Interphase	S-Phase
place during				-	
Rough endoplasmic	RNA	Ribosome	Protein	DNA	Ribosom
reticulum is embedded					e
with					
Hershey and Chase	T2-Phages	Salmonella	Amoebae	Bacterium	T2-
experiment on					Phages
RNA has the following	ATG	AUG	APG	ATT	AUG
bases					
Chromatin composed of	Nucleic acids and	Nucleic acids only	Proteins only	RNA only	Nucleic
-	proteins			-	acids and
					proteins
Nucleosome assembly	initiated by	spontaneous	inhibited by	induced	
is	nucleoplasmin		topoisomerase		spontane
					ous
Polytene chromosomes	gene inversion	Gene conversion	DNA	gene deletion	DNA
are produced by	-		amplification	-	amplifica
					tion

In DNA double helix, the two DNA chains are held together by bonds between the base pairs	Covalent	Hydrogen	Ionic	Hydrophobic	Hydroge n
Smaller blocks occur at the end of chromosome arm is called	centromere	Telomere	blastomeres	blastocyst	Telomere
Proteins that can unwind a DNA helix are	Helix destabilizing proteins	Melting proteins	Helix destabilizing proteins and Melting proteins	Single strand binding proteins	Helix destabiliz ing proteins and Melting proteins
The length of the DNA segment present in the nucleosome core particle is	140 bp	200bp	166 bp	114 bp	140 bp
The molecular chaperone which causes the nucleosome assembly is	nucleoplasmin	histone	Hu protein	Ubiquitin	histone
The base pair between adenine and thymine forms hydrogen bonds	2	1	3	4	. 2
Bacterial transformation is	Griffith effect	Averys effect	Hershey effect	Joule effect	Griffith effect
The process DNA duplication is	Replication	Transcription	Translation	Reverse transcription	Replicati on
The process of making copy of RNA from DNA is	Replication	Transcription	Translation	Reverse transcription	Transcrip tion
The process of synthesizing protein from RNA	Replication	Transcription	Translation	Reverse transcription	Translati on
The process of making copy of DNA from RNA is	Replication	Transcription	Translation	Reverse transcription	Reverse transcript ion
Satellite DNA is rich inbase sequence		GC	GCAT	GT	AT
Important chaecteristic of satellite DNA is, they contain	repetitive base sequence	GC rich	Unique sequence	solitary sequence	repetitive base sequence

Chicken lysozyme gene is a good example of	Single copy DNA	Moderatively Repetitive DNA	Simple sequence DNA d.	highly Repetitive DNA	Single copy DNA
In linker DNA, which histone is involved	H1	H2	Н3	H4	H1
The sister chromatids will be attached by	a centromere	a telomere	a centriole	golgi bodies	a centrome re
is the division of the cytoplasm, separating the organelles and other cellular components.	Mitosis	Meiosis	Cytokinesis	Proface	Cytokine sis
Chromatin condenses into chromosomes, the nucleolus dissolves, nuclear membrane is disassembled, and the spindle apparatus forms in	Prophase	Metaphase	Anaphase	Telophase	Prophase
Simple non-nucleated cells are	Prokaryotic cells	Eukaryotic cells	Stem cells	Blood cells	Prokaryot ic cells
Chromosomes are seen in the equatorial plane in	Prophase	Metaphase	Anaphase	Telophase	Metaphas e
An interphase between nucleus and cytoplasm is	Nuclear envelop	Nuclear membrane	Nuclear pores	Perinuclear space	Perinucle ar space
The cell cycle consists of	two distinct phases	three distinct phases	four distinct phases	five distinct phases	four distinct phases
Sucide bags are otherwise called	Ribosomes	Lysosomes	Golgi apparatus	Endoplasmic reticulum	Lysosom es
The Golgi apparatus important for	Protein synthesis	DNA synthesis	RNA synthesis	Packaging and secretion of proteins	Packagin g and secretion of proteins
Rough endoplasmic reticulum is embedded with	RNA	Ribosome	Protein	DNA	Ribosom e
In rolling circle replication single stranded viral DNA otherwise called as	Positive strand	Negative strand	Replicative strand	Rolling strand	Positive strand
Which of the following catalize the formation of negative coils during DNA replication?	helicases	gyrases	primosome	polymerase	gyrases

In Hershey and Chase	DNA	Proteins	Cytoplasm	RNA	Proteins
experiment S ³⁵ is					
incorporated in					
formation of					

The DNA replication is semi conservative	Meselson &Stahl	Okazaki	Lehniger	Lorunberg	Meselson &Stahl
was demonstrated by					
The process DNA duplication is	Replication	Transcription	Translation	Transversion	Replication
The seperation of two DNA strands for	DNA polymerase	DNA helicase	DNA	DNA ligase	DNA helicase
replication is brought by	III		polymerase I		
Which of the following	(5')TCTGACCAG((5')CTCATTGA	(5')GAGTCAAC	(5')GACCAGTCT	(5')GACCAGTCT(3
deoxyoligonucleotides will hybridize with a	3')	G(3')	T(3')	(3')	')
DNA containing the sequence					
(5')AGACTGGTC(3')?					
Negative supercoils are removed by	Topoisomerase I	gyrase	helicase	rep protein	Topoisomerase I
The term Okazaki fragments refers to the	discontinuously	discontinuously	DNA fragments	DNA fragment	discontinuously
	synthesized DNA	synthesized DNA	formed by the	from pol I	synthesized DNA
	pieces on the	pieces on the	action of DNA		pieces on the
	lagging strand	leading strand	ligase		lagging strand
		_	-		
The synthesis of new DNA (replication) in		DNA	DNA polymerase	DNA polymerase	DNA polymerase
prokaryotes is catalysed by	DNATopoisomeras	polymerase I	II	III	III
	e				
Okazaki fragements are initiated with	DNA primer	RNA primer	DNA template	RNA template	RNA primer
Semiconservative mechnism of replication	Crick	Watson	Watson & Crick	Benjamin	Watson & Crick
was proposed by				&Franklin	
Eukaryotic cells have kinds of RNA	1	2	3	4	3
polymerase					
The lagging strand is characterized by	a constant supply	discontinuous	replication of	continuous	continuous
except	of RNA primers	synthesis of new	DNA in 5'-3'	synthesis of new	synthesis of new
		strand	direction	strand	strand
Replication is characterized by the	semiconservative	bi-directional		dependent on	dependent on DNA
following except it is			semidiscontinuou	DNA primer	primer
			S	-	
The process linking two circular DNA	decatenation	catenation	annealing	nick translation	catenation
molecules to form a chain is called					

The rep protein moves along the	leading strand	lagging strand	leading strand	lagging strand	lagging strand
	template in 3'-5'	template in 3'-5'	template in 5'-3'	template in 5'-3'	template in 3'-5'
	direction	direction	direction	direction	direction
Ori C is rich insequence	GC	AT	ATGC	GATC	AT
The coding strand of DNA is	the plus strand	the minus strand	the antisense strand	template strand	the plus strand
A replicon is	region with DNA origin	regulatory gene	Inverted repeats	tandom repeats	region with DNA origin
The DNA replication is discontinuous was proved by	Messelson-Stahl	Reigi Okazaki	Albert Lehninger	Arthur Kornberg	Reigi Okazaki
Watson and Crick elucidated ds DNA structure by using	NMR spectroscopy	X-ray diffraction	Circular dichroism	IR and Raman spectroscopy	X-ray diffraction
. The enzyme primarily involved in DNA repair in bacteria is	DNA pol I	DNA pol II	DNA pol III	DNA pol a	DNA pol I
In bacteria, which enzyme unwinds, binds ss DNA, denatures ds DNA and matches the ss DNA with complementary denatured DNA ?	Rec A	Uvr ABC	Uvr D	Dna B	Dna B
The melting temperature of DNA is	Directly propotional to A-T content	Directly propotional to G- C content	Not related to base composition	Directly propotional to DNA length.	Directly propotional to G-C content
Genetic information is perpetuated by	repair	transcription	translation	replication	replication
The catenated circules in E. Coli are separated by	Topoisomerase I	Topoisomerase II	Topoisomerase III	Topoisomerase IV	Topoisomerase IV
E. Coli DNA ligase utilizes	NAD+	ATP	GTP	NADH	NAD+
is an E.Coli protein that melts duplex DNA	DNa A	DNa B	DNa C	DNa D	DNa A
The Klenow fragment exhibits the activity of	5'-3' exonuclease	polymerase and 3'-5' exo nuclease	polymerase and 5'-3' exo nuclease	an endo nuclease	polymerase and 3'- 5' exo nuclease

For the polymerase I activity	primer is not	primer with	Primer with free	Primer with free 5'	Primer with free 3'-
	required	either free 3' OH	3'-OH is required	OH is required	OH is required
		or 5' – P is	_	_	_
		required			
DNA polymerase I has all the activities	3'- 5' exonnclease	5'-3'	endonuclease	Helicase	Helicase
except	activity	exonnclease	activity		
		activity			
Among the following which is a substrate for DNA polymerase I?	5' monophosphate	5' diphosphate	5'triphosphate	deoxy ribose	5'triphosphate
In DNA replication helix unwinding is accomplished by enzymes called	helicases	gyrases	topoisomerases	polymerases	helicases
The term processivity with reference to	the rate at which	the number of	the rate at which	fidelity of DNA	the number of
DNA polymerase refers to	polymerization	nucleotides	the enzyme	replication	nucleotides added
	occurs	added before the	dissociates from		before the enzyme
		enzyme	the template		dissociates from the
		dissociates from			template
		the template			
The initiation of a new round of replication	dna C	dna B	dna A	pol A	dna C
in E.coli is regulated by the gene					
RNA primers are removed by	DNA polymerase I	DNA	DNA	topoisomearse	DNA polymerase I
		polymerase II	polymerase III		
E.coli DNA replicase is	Pol III	Pol II	pol I	ligase	Pol III
The two replicating forks travel in opposite	termini	replication fork	intron	exon	termini
directions until they reach either end of that					
unit, the two end points are called					
dna B code for the enzyme	helicase	primse	polymerase	ligase	helicase
The presence of following four deoxy	dATP, dUTP,	dATP, dGTP,	dATP, dGTP,	dATP, dGTP,	dATP, dGTP,
ribonucleoside triphosphate is essential for	dCTP, dTTP	dUTP, dTTP	dCTP, dTTP	dCTP, dUTP	dCTP, dTTP
replication					
The synthesis of new DNA (replication) in	DNA polymerase I	DNA	DNA polymerase	DNA topo	DNA polymerase
prokaryotes is catalysed by		polymerase II	III	isomerase	III

Single strand binding protein binds to single	-	. to repair base	to initiate	to prevent	to prevent
strand DNA	replication	pairs	transcription	reformation of	reformation of
				duplex state.	duplex state.
Which of the following possesses both 5'-3'	Kornberg enzyme	DNA polymerase	Taq DNA	DNA polymerase I	Kornberg enzyme
and 3'-5' exonuclease activity?		III	polymerase		
In E.coli, which enzyme synthesizes the	DnaA	DnaB	DnaC	DnaG	DnaG
RNA primer for Okazaki fragments?					
DNA gyrase is inhibited by	tetracycline	Nalidixic acid	Streptomycin	cephalosporin	Nalidixic acid
DNA replication rates in prokaryotes are	10 bases per	100 bases per	1000 bases per	10,000 bases per	1000 bases per
approximately of the order of	second	second	second	second	second
During which of the following process a new	Trasformation	Transcription	Translation	Replication	Replication
copy of a DNA molecule is precisely					
synthesized?					
Which of the following repairs nicked DNA	Helicase	DNA gyrase	Topoisomerases	DNA ligase	DNA ligase
by forming a phosphodiester bond between					
adjacent nucleotides?					
The scientist involved in discovery of DNA	Hershey and Chase	Griffith and	Avery, MacLeod	Waston and Crick	Avery, MacLeod
as chemical basis of heredity were.		Avery	and McCarty		and McCarty
Which of the following help to hold the	primase	ligase	DNA polymerase	single strand	single strand
DNA strands apart while they are being				binding proteins	binding proteins
replicated?					
What is the role of DNA ligase in the	synthesize RNA	catalyze the	join Okazaki	unwind the	join Okazaki
elongation of the lagging strand during DNA	nucleotides to	lengthenin	fragments	parental double	fragments together
replication?	make a primer	g of telomeres	together	helix	
Which of the following covalently connects	helicase	DNA polymerase	ligase	DNA polymerase I	ligase
segments of DNA?		III			
What is the function of helicase?	It forms bonds	It adds new	It forms the DNA	It separates DNA	It separates DNA
	between DNA	nucleotides to the	helix	strands	strands
	nucleotides	DNA helix.			
Nucleosome assembly is	initiated by	spontaneous	inhibited by	induced	spontaneous
	nucleoplasmin	_	topoisomerase		_
The coiling that cannot be separated except	supercoiling	negative super	plectonemic	anti parellel	plectonemic
by unwinding is called		coiling	coiling	coiling	coiling

If replication was completely conservative	one heavy and one	both heavy	both light strands	Mixed strands	one heavy and one
then	light strand would	strands would be	would be seen		light strand would
	be seen	seen			be seen
Specific chromosomal locations from where	locus	origin of	loci	chromosomal arms	origin of replication
replication begins are called		replication			
All biological DNA synthesis occurs from	3 end to 5 end	5 end to 3 end	both A and B	5' end to 5' end	5' end to 3' end
the					
Transition occurs when purine is substituted	pyrimidine	pyrol	purine	phenazine	purine
with					
Change in nucleotide sequence of a short	replication	recombination	repair	mutation	mutation
region of genome is called					
When DNA opens up y shaped structures are	replication forks	single stands	parent strand	double strand	replication forks
called					
All newly synthesized polynucleotide strands	RNA polymerase	DNA polymerase	gyrase	helicase	RNA polymerase
must be initiated by					

5' cap in mRNA is done by	ATP	GTP	СТР	UTP	GTP
Large protein complex that	replisome	primosome	spliceosome	Chromosome	
involved in intron removal is					spliceosome
One of the following is distinctive	poly A tail	poly T tail	polu U tail	poly G tail	poly A tail
Structure of eukarytic mRNA					
by	DNA Pol I	RNA Pol I	RNA Pol II	RNA Pol III	RNA Pol II
The strand of DNA that directs the synthesis of mRNA is	sense strand	leading strand	antisense strand	lagging strand	sense strand
Primase initiates all excxept	leading strand synthesis	replication	Okazaki fragments	translation	translation
Positive superhelicity generated during DNA replication is removed	DNA polymerase	DNA isomerase	DNA gyrase	DNA helicase	DNA gyrase
The problem of supercoils during	DNA	DNA ligases	DNA	DNA helicases	DNA
DNA replication is overcome by a	topoisomerases		polymerases		topoisomerases
group of enzymes called					
The DNA element enhancer	can act only in cis	. can act only in	can act in cis or	can act at medial	can act only in cis
		trans	in trans		
Telomerase is a	primase	polymerase	reverse	helicase	reverse transcriptase
		4	transcriptase		
For initiationsubunit of RNA pol required	alpha	beta	gamma	sigma	sigma
Pribnow box consists of the	TATAAT	TATATA	TTAAAT	TAATAT	TATAAT
following 6 nucleotide bases					
Rho protein is involved in	transcription	transcription	transcription	. transcription	transcription
	initiation	elongation	termination at	termination at all	termination at some
			some terminators	terminators	terminators
5SrRNA are synthesized by RNA	I	II	III	IV	III
polymerase					
Transfer RNA has all except	Clover leaf	Anticodon arm	Minor bases in	proteins	proteins
	structure		addition to		
			normal bases A,		
			U, G and C		

snRNAs are involved in the	Transcription	Splicing of		splicing of tRNA	Splicing of mRNA
eukaryotic		mRNA	acids		-
Rho factor is a	Dimer	Trimer	hexamer	Tetramer	hexamer
RNA self splicing was discovered	Watson & Crick	McClintock	Sanger	Thomas Cech	Thomas Cech
by					
Transcription charecterised by all	one strand of DNA	synthesis of	Synthesize	Participation of	Synthesize proteins
except	serve as template	RNA	proteins	RNA polymerase	
Termination of transcription is brought about by	sigma factor	RNA polymerase	rho factor	TATA box	rho factor
In eukaryotic cells, the synthesis of	RNA polymerase I	DNA polymorogo	DNA polymorese	DNA polymorese	RNA polymerase II
	KINA porymerase i			DNA porymerase	KINA polymerase m
mRNA is carried out by		Π			, •
Rho factor is	an enzyme	accessory	rich in U residues	rich in G-C with a	accessory protein
		protein		palindromic	
				sequence.	
The protein that bind to the TATA	coregulators	coactivators	. enhancers	Transcriptional	Transcriptional
box in the promoter region are				factors	factors
Who demonstrated that poly U	Nirenberg and	. Nirenberg and	Nirenberg and	H. Gobind	Nirenberg and
stimulate the synthesis of poly	Leder	Ochoa	Mathai	Khorana	Mathai
Shine dalgarno sequence is rich in	purines	. pyrimidines	. only A	only T	purines
The enzyme responsible for	Rnase III	Rnase H	Rnase P	Rnase F	Rnase III
processing the rRNAs is					
Hirpin loop like structure in	Initiation	Elongation	termination	Initiation&elongati	termination
transcription is related to				on	
Proceesing of tRNA involves	CCA to the 3' end	CCA to the 5'	7-methyl G to	poly A to the 3'	CCA to the 3' end
adding		end	the 3' end	end	
A tRNA molecule must be able to	recognize a codon	recognize an	distinguish one	recognize DNA	recognize a codon
	6	anticodon	amino acid from	molecules	6
			another		
The following are associated with	-35 sequence	pribnow box	promoter	spacer	spacer
transcription except	20 bequence	Priorio a cox	Promotor	Spacer	Space
What is the characteristic form	Linear structure	circular form	lariat shaped	theta structure	lariat shaped
introns have after being cut from a			harran shaped		
muons nave aller being cut nom a	1		1	1	

During the course of transcriptuon or replication for the addition each	phosphate	pyrophosphate	nitrogenous base	nucleotide	pyrophosphate
nucleotide to the growing chain the following moiety is released					
The RNA polymerase of E.coli	2a,2b and one	2a 2b and one	2a 1b,1b and one		2a 1b,1b and one
complex holoenzyme composed of 5 polypeptide subunits	sigma factor	rho factor	sigma factor	rho factor	sigma factor
RNA polymerase utilizes the	ATP, GTP, CTP	ATP, GTP, CTP	ATP, CTP, UTP	CTP, UTP	ATP, GTP, CTP
following RNA triphosphates for the formation of RNA	and TTP	and UTP	and UTP		and UTP
tRNA specifically reacts with	mRNA	nuclear RNA	amino acyl	ribosomal RNA	amino acyl
			adenylates		adenylates
Pseudo uridine is found in	mRNA	snRNA	rRNA	tRNA	tRNA
mRNAs are synthysized by RNA polymerase	. I	Π	III	I & II	Π
Mature mRNAs has the following	5'Cap	Poly A tail	Exon	Intron	Intron
properties except Rho factor catalyses the unwinding	Hydrolysis of	RNA-RNA	Microsco alona	Microtos alona	Understroig of ATD
of RNA-DNA double helices and	ATP	double helices	Migrates along the RNA in 3'-5'	Migrates along the RNA in 5'-3'	Hydrolysis of ATP
		double hences	direction	direction	
Prokaryotic transcription initiation is inhibited by	. Rifamycin	Rifampicin	ricin	abrin	Rifampicin
Non template strand is otherwise called as	template strand	coding strand	non coding strand	sense strand	coding strand
activates transcription	CRP	GMP	RNase	ribozyme	CRP
The region on t RNA recognizes	codon loop	anti-codon loop	. P site	. A site	anti-codon loop
the appropriate base sequence m					
RNA is called					
Enhancers are entry point on DNA for RNA polymerase	Ι	Π	Ш	I and III	П
Promoters are recognized by	alpha sub unit	gamma subunit	beta subunit	sigma subunit	sigma subunit
Eukaryotic mRNA that specifies	Addition of 5' cap	removal of	addition of 3'	6	removal of introns
proteins is modified before		exons	poly A tail		

The 5' terminus of a eukaryotic	2-amino purine	5 bromo uracil	7 methyl	formyl	7 methyl guanosine
mRNA molecule is			guanosine	methionine	
. When a number of genes are transcribed as one mRNA, the mRNA is said to be	multimeric	. polymeric	polycistronic	polyclonal	polycistronic
Transposase is encoded by	IS element	bacterial trnasposon	. rRNA	. tRNA	bacterial trnasposon
Premature chain termination in prokaryotes and eukaryotes is	ricin	abrin	erythromycin	. puromycin	ricin
Rho dependant termination sites	only U	. Only GC	Both U and GC	. AT	Both U and GC
Internal promoters are present in the gene sequence of	18SrRNA gene	28SrRNA genes	mRNA genes	t RNA	t RNA
The hexa nucleotide AAU AAA provides the signal for	Capping	Generation of eukaryotic 3'- poly A tail	Initiation of translation	Peptide bond formation	Generation of eukaryotic 3'- poly A tail
RNA polymerases present in eukaryotes are	RNA pol I	RNA pol II	RNA pol III	RNA pol I,II and III	RNA pol I,II and III
The mature 5'end of tRNA in E.coli formed as a result of endonucleolytic cleavage by	RNase P	RNAse D	RNase M5	RNase M13	RNase P
The enzyme responsible for reverse transcription	DNA dependent DNA polymerase	1	RNA dependent DNA polymerase	RNA dependent RNA polymerase	RNA dependent DNA polymerase
DNA Polymerase I has	5'-3' Exonuclease activity	Proof reading activity	Polymerising activity	5'-3' Endonuclease activity	Polymerising activity
An E.Coli strain lacking DNA polymerase I would be deficient in	Repair	Splicing	Methylation	Transcripition	Repair
Single standered circular DNA is present in	j/174 bacteriophages	all viruses	Animal viruses	Plant viruses	φ/174 bacteriophages
In the classical model of transcriptional control described by Jacob and Monod, a repressor	Enhancer	AUG sequence	Operator	Ribosome binding site	Operator

Translation refers to	DNA from RNA	RNA from DNA	protein from DNA	protein from RNA	protein from RNA
The chain initiating amino acid in protein biosynthesis in E.Coli is	Methionine	. Cysteine	N-formyl methionine	. Arginine	N-formyl methionine
are the factories for protein synthesis	mitochondria	nucleus	ribosomes	cytoplasm	ribosomes
The initiating codon in protein synthesis	UAA	. UAG	UGA	AUG	AUG
Genetic code is the dictionary of the	A, G, C and T	A, G, C and U	. A, G, T and U	. A, G, T and T	A, G, C and U
following nucleotide bases					
The Wobble hypothesis accounts for the	triple nature of codon	degeneracy of codon	universality of the codon	doublet nature of the codon	degeneracy of codon
Which of the following amino acids is specified by only a single codon?	. Glutamine	. tryptophan	asparagine	isoleucine	isoleucine
The enzyme responsible the cleavage of the bond connecting amino acid and tRNA is	amino acyl-tRNA synthetase	peptidyl transferase	aminoacyl transferase	tRNA deacylase	tRNA deacylase
Catalytic RNA is	Abruma	rihoruma	anguma	protein	rihoruma
	Abzyme	ribozyme	enzyme	protein	ribozyme
are proteins that block the synthesis of RNA at gene level	Initiators	Repressors	stimulators	retarders	Repressors
tRNA, 5S RNA, Special RNA s are synthesised by RNA Pol	I	Ш	ш	IV	Ш
Which of the following antibiotics is an inhibitor of protein synthesis	Chloramphenicol	Penicillin	Bacitracin	Cephalosporin	Chloramphenicol
Coupled transcription and translation occurs in	only in eucaryoptes	only in prokaryotes	both in prokaryotes and in eukaryotes	neither in prokaryotes nor in eukaryotes	only in prokaryotes
Which of the following are termination codons	UAG, UAA, UGA	UUA, AUC, GUC	. UAG, AGG,UGA	. UAG, GAA, GUC.	UAG, UAA, UGA
Nucleic acid concerned with the protein synthesis is	DNA	mRNA	. ribosomal RNA	tRNA	. ribosomal RNA
One of the following compound competitively inhibits the enzyme peptidyl transferase and interfere with elongation of peptide chain	Streptomycin	Puromycin	Chloramphenicol	Terracycline	Chloramphenicol
The stop codon UAA is otherwise called as	Amber	Ochre	opal	none	Ochre
The binding of tRNA ^{fmet} to the P site is inhibited by	Streptomycin	. Chloramphenicol	Erythromycin	Lincomycin	Streptomycin
Translocase is an enzyme required in the process of	.DNA replication	RNA synthesis	Initiation of protein synthesis	Elongation of peptides	Elongation of peptides
Which one of the following bases if present in the third position of an anticodon cannot form wobbl base	G	С	U	A	Α
How many polypeptide chains can be formed simultaneously by a given ribosome?	one	about a dozen	up to 30	variable depending on mRNA	one
Diphtheria toxin inhibits elongation step in translation through	Glycosylation	Methylation	ADP- ribosylation	Ubiquitinylation	ADP- ribosylation
Selection of translational initiation site is carried out by	Base pairing between mRNA and 16rRNA	initiation codon	termination codon	promoter	Base pairing between mRNA and 16rRNA
Principle function of RNA pol II in Eukaryote is synthesis of	mRNA	tRNA	rRNA	tRNA	mRNA
Which of the following inhibits the peptidyl transferase activity of the eukaryotic 60S ribosomal subunit	Chloroamphenical	Cycloheximide	Erythromycin	Puromycin	Cycloheximide
Large subunit of ribosome is present ineukaryotes	60s	. 40s	30s	80s	60s

In protein synthesis aminoacids are joined	peptide	phosphodiester	disulfide	phosphodiester and disulfide	peptide
by bond	peptide	phosphoulester	disuffice		popula
Which of the following Binds to 50S subunit and inhibits translocation	Chloroamphenical	Cycloheximide	Erythromycin	Puromycin	Erythromycin
The codon that terminates protein synthesis is	UGG	UGA	UGC	UGU	UGA
Shine Dalgarno sequence is	rich in pyrimindine	rich in purine	centered 100 nucleotide upstream from the start codon	centered 100 nucleotide downstream from the start codon	rich in purine
The following are the termination or non sense codons except	AUG	UAA	UAG	UGA	AUG
The specific information required for the sequence amino acids in protein is located on	rRNA	mRNA	tRNA	hnRNA	mRNA
The largest subunit of ribosomal RNA in prokaryotes	308	50S	60S	40S	508
Which one of the following post translational modifications occurs in clotting factors?	hydroxylation	methylation	carboxylation	phosphorylation	carboxylation
The antibiotic cycloheximide which	blocks chain elongation during protein synthesis	blocks glycosylation of proteins	resembles aminoacyl tRNA	inhibits the formation of peptide bond.	inhibits the formation of peptide bond.
Post dimer initiation is responsible for	recombination repair	SOS repair	excision repair	photo reactivation	recombination repair
The association of 50S and 30Ssubunit is prevented by	IF 1	IF 2	IF 3	EF2	IF 3
Wobble hypothesis explained by one of the following characteristics of the genetic code	Universality	Specificity	non overlapping	degenerate	degenerate
The codon (of mRNA) and anticodon (of tRNA) recognize each other by pairing	5'-3' of mRNA with 5'-3' of tRNA	3'-5' of mRNA with 3'-5' of tRNA	5'-3' of mRNA with 3'-5' of tRNA	3'-5' of mRNA with 5'-3' of tRNA	5'-3' of mRNA with 3'-5' of tRNA
Proteins are synthesized in direction	Amino to carboxyl	Carboxyl to amino	Both amino and carboxyl direction	other groups like phosphoric acid	Amino to carboxyl direction
Post translational modifications involve the following covalent changes	Synthesis mRNA	hydroxylation	carboxylation	Phospho diester linkage formation	Phospho diester linkage formation
Wobble hypothesis is characterized by recognition of	a single codon by a single tRNA	more than one codon by asingle tRNA	a single codon by more than one tRNA	more than one codon by more tRNA	more than one codon by asingle tRNA
Rifampicin inhibit Protein synthesis by binding to	beta subunit of RNA pol	delta subunit of RNA pol	gamma subunit of RNA pol	alpha subunit of RNA pol	beta subunit of RNA pol
In prokaryotes protein synthesis is carried out by ribosomes of	80s,30s, and 50s	80s,40s, and 50s	70s,30s, and 50s	70s,40s, and 60s	70s,30s, and 50s
Premature termination of poly peptide is caused by	Erythromycin	Puromycin	Streptomycin	Ricin	Puromycin
Which of the following is not a necessary component of translation?	Anticodon	mRNA	Ligase	Amino acid	Ligase
Eukaryotic mRNA binding to the ribosomes is facilitated by	the Shine Dalgarno sequence	the 7-methyl guanosine cap	tRNA	poly A tail	the 7-methyl guanosine cap
The structure in a bacterium that indicates an active site for protein synthesis is	a chromosome.	a cell membrane,	a flagellum.	a polysome.	a polysome.
In the presence of high levels of tryptophan	attenuator allows transcription of trp structural genes	attenuator propogates transcription	attenuator terminates transcription	attenuator terminates translation	attenuator terminates transcription

Which of the following occur in the presence of glucose?	lac Z gene expression is increased		Binding of CAP-cAMP complex to the promoter area decreases		Binding of CAP-cAMP complex to the promoter area decreases
The lac operon is translated into proteins.	1	2	3	4	3
In the presence of tryptophan in the cell, the repressor bound to	tryptophan	DNA	DNA and tryptophan	neither tryptophan nor DNA	DNA and tryptophan
Catabolite activating protein exerts control on transcription of the lac operon genes.	positive	negative	may be positive or negative	positron	positive
DNA is methylated at	AC sequences	TATA sequences	CAAT sequences	CG sequences	CG sequences
Which type of ribonucleic acid combines with proteins to form a ribosome?	mRNA	tRNA	sRNA	rRNA	rRNA
How many bases are needed to code for two amino acids?	2	3	6	12	6
What does the operon model attempt to explain?	the coordinated control of gene expression in bacteria		how genes move between homologous regions of DNA	the mechanism of viral attachment to a host cell	the coordinated control of gene expression in bacteria
A lack of this nonprotein molecule would result in the inability of the cell to "turn off" genes:	inducer	promoter	repressor	corepressor	corepressor
The lactose operon is likely to be transcribed when	there is more glucose in the cell than lactose.	the cyclic AMP levels are low.	there is glucose but no lactose in the cell.	the cyclic AMP and lactose levels are both high within the cell.	the cyclic AMP and lactose levels are both high within the cell.
During translation, chain elongation continues until what happens?	All tRNAs are empty	The polypeptide is long enough	A stop codon is encountered.	The ribosomes run off the end of mRNA.	A stop codon is encountered.

The process of DNA repair in which enzymes cut out a damaged section of DNA and other enzymes then repair the resulting break is known as	excision repair	recombination repair	SOS repair	Photo reactivation	excision repair
Removal of thymine dimer after replication process is done by	recombination repair	excision repair	light repair	photo reactivation	recombination repair
Excision repair is carried by	Uvr ABC endonuclese	AP endonuclease	Methyl transferase	Glycosylase	Glycosylase
In the following statement all are related to photoreactivation repair except	an enzymatic cleavage of thiamine dimers	activated by visible light	uses light energy to cleave C-C bonds of cyclobutyl rings	removal of short segment of DNA	removal of short segment of DNA
The enzyme responsible for the replication of mitochondrial DNA	DNA polymerase α	DNA polymerase β	DNA polymerase γ	DNA polymerase δ	DNA polymerase γ
5-bromo uracil damages DNA. It is an anolog of	Uracil	Thymine	Cytosin	Guanine	Thymine
The type of point mutation which could be detected only after nucleic acid sequencing is known as	silent mutation	missense mutation	nonsense mutation	. Transversion mutation	silent mutation
Which of the following enzyme selectively degrades single stranded nucleic acid	DNA ligase	. DNA polymerase	Ribonuclease	S1 nuclease	S1 nuclease
Pyrimidine dimers are mended by all except	excision repair	photo lyase	recombination repair	Mismatch repair	Mismatch repair
Altered bases in DNA are removed by	Photolyase	Glycosylase	Apendo nuclease	UvrABC endonulease	Glycosylase
Hot spots are sites containing Which of the following have proof reading activity	MeC Rep protein	MeG polymerase I	MeA polymerase II	MeA polymerase III	MeC polymerase I

Which of the following thymine	excision repair	recombination	SOS repair	Photo reactivation	Photo reactivation
dimer repair pathway is		repair			
dependent on light?					
Daughter strand repair is the	SOS repair	excision repair	recombination	photo reactivation	recombination repair
other name for			repair		
Dna A protein recognize and	49 bp repeats in	13 bp AT rich	49 bp repeats in	RNA polymerase	4 9 bp repeats in
binds to	ter C	segment	Ori C		Ori C
Discrimination of methylation	mismatch repair	excision repair	recombination	photo reactivation	mismatch repair
in GATC sequence of parent			repair		
and daughter strand is observed					
Rec A protein can bind to	ssDNA	ds DNA	RNA	protein	ssDNA
Chemical adducts are DNA	having chemically	having unusal	rich in AT	rich in GC	having chemically
region	modified bases	bases			modified bases
The proof reading activity of the	DNA helicase	DNA	DNA	DNA polymerase	DNA polymerase I
newly synthesized DNA is		polymerase I	polymerase II	III	
present on the enzyme					
The type of point mutation	silent mutation	missense	nonsense	. Transversion	silent mutation
which could be detected only		mutation	mutation	mutation	
after nucleic acid sequencing is					
known as					
Common lesions found in DNA	Pyrimidine dimers	Single strand	Base deletion	Purine dimers.	Pyrimidine dimers
after exposure to ultraviolet light		breaks			
Genetic recombination in	Lederberg and	Hershey and	Avery and	Griffith	Lederberg and
bacteria discovered by	tautum	chase	Mccarthy		tautum
Exchange of a single nucleotide	transition	transversion	recombination	translation	transition
$(A \leftrightarrow G)$ or $(C \leftrightarrow T)$.is					
Point mutation which code for	missense mutation	nonsense	silent mutation	inseration	missense mutation
the differnt amino acid is		mutation			
Point mutation which code for a	missense mutation	nonsense	Silent mutation	inseration	nonsense mutation
stop and truncate the protein is		mutation			

Mutation occur in non coding	missense mutation	nonsense	Silent mutation	inseration	Silent mutation
region is		mutation			
The biochemical mechanism tolerance of the damage is	SOS repair	Excision repair	Recombination	reverse mutation	SOS repair
Includes a bypass system that allows DNA chain growth across damaged segment at the cost of fidelity of replication	SOS repair	Excision repair	Recombination Excision repair	Incision repair	SOS repair
The drug chloramphenical blocks	cellwall formation	transcription	translation termination release factors	polypeptide chain elongation.	polypeptide chain elongation.
AT repeats are the examples of	insertion	deletion	transversion	translocation	insertion
A base is changed by the repositioning of a hydrogen atom.	tautomerism	depurination	deamination	transition	transition
Loss of a purine base (A or G)	tautomerism	depurination	deamination	transition	depurination
A purine changes to another purine, or a pyrimidine to a pyrimidine	tautomerism	depurination	deamination	transition	transition
which of the following is an alkylating agent	NTG	BU	NH2OH	HA	NTG
Common lesions found in DNA after exposure to ultraviolet light	Pyrimidine dimers	Single strand breaks	Base deletion	Purine dimers.	Pyrimidine dimers
During mutation substitution of Adenine by Guanine is known as	Transition	Transversion	Deletion	Transpositions	Transition
Mutation leads to the death of all	Lethal	Point	Frame shift	Missense	Lethal
Which of the following leads to frame shift mutation	base analog	ionization	uv radiation	5 BU	uv radiation
Which one is an mutagen	EMS	EES	NTG	HA	EMS

Which of the following is not require for plasmid to select recombination?	ori	marker site	restriction enzyme site	recombination site	recombination site
Transposons are	Regulatory elements	Jumping genes	Enzymes	restriction enzymes	Jumping genes
Transposable elements were discovered by	Mendel	Darwin	McClintock	Tomizerwa	McClintock
One of the following is not a kind of mobile genetic elements?	Operon	Plasmid	Transposon	Bacteriophage.	Operon
The insertion sites used repeatedly are called	Cols spots	Hot spots	Random spots	Unique spots	Hot spots
The termini of each Is elements have	Inverted repeat sequence	Initiation repeat sequence	Contrast repeat sequence	Plasmid repeat sequence	Inverted repeat sequence
LTR strands for	Long terminal repeats	Left terminal repeats	Large terminal repeats	L terminal repeats	Long terminal repeats
The genetic organization of the Ty elements resembles that of the eukaryotic	Retro viruses	Mosaic viruses	Sendai viruses	Animal viruses	Retro viruses
The genes present in a provirus's is	Gap gene	Pol gene	Env gene	Tus gene	Gap gene
The yeast Ty elements are sometimes called as	Simple transposons	Complex transposons	Retro transposons	Co- transposons	Simple transposons
The Holliday intermediate is also known as the	α-form	β form	μ- form	form -א	form -א
Holliday model of recombination is observed in which stage of cell cycle	DNA Synthesis	G1	G2	Meiosis	Meiosis
Repressor binding site on DNA is	Activator	Operator	enhancer	stimulators.	Operator
House keeping gene is the gene that expressed	rarely	constantly	over expressed	under expressed	constantly
Lac repressor is a	Monomer	Dimer	Trimer	Tetramer	Tetramer

Antisnse RNA is related	PCR	RNAi	Southern blot	Western blot	RNAi
tRNA acts as	intergrator	Adaptor	regulator	promoter	Adaptor
molcule					
Double strand RNA is degraded	endouclease	protease	RNase	Dnase	RNase
by					
Site specific recombination require the recognition of unique nucleotide sequences in both DNA molecules by enzymes	recombinase	Helicase	Isomerase	ligase	recombinase
Recombination in Meselson- Radding model is initiated by	ss nicks	ds nicks	exchange process	mispairing	ss nicks
RNA polymerase binds to DNA at	activator	Operator	Promoter	repressor	Promoter