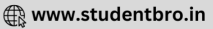
BIOTECHNOLOGY PRINCIPLES AND PROCESSES

1.	First hormone prepared	by genetic engineering is:		
	a) Insulin	b) Oxytocin	c) Adrenaline	d) Somatotropin
2.	Retroviruses in animals i	ncluding humans are able t	o change normal cells into	
	a) Germ cell	b) Cancerous cells	c) Cosmid	d) Vector
3.	The restriction enzyme r	esponsible for the cleavage	of following sequence is	
	5' - G - T - C - G - A -	· C − 3′		
	3' - C - A - G - C - T -	G - 5'		
	a) <i>Alu</i> I	b) Bam HI	c) Hind II	d) <i>Eco</i> RI
4.	pBR322 was the first arti	ficial cloning vector develo	ped inA byB and	C from <i>E. coli</i> plasmid.
	Here A, B and C can be			
	a) A-1976, B-Boliver, C-R	lodriquez	b) A-1975, B-Tiselius, C-F	Rodriquez
	c) A-1977, B-Boliver, C-R	todriquez	d) A-1978, B-HO Smith, C	-KW Wileox
5.	Transfer of any gene into	a completely different org	anism can be done through	
	a) Genetic engineering	b) Tissue culture	c) Transformation	d) None of these
6.	An environmental agent	that triggers transcription f	from an operon is a:	
	a) Depressor	b) Inducer	c) Regulator	d) Controlling element
7.	Recombinant DNA have i	[1] 발생 보다 다음 마음 보다 보는 것이 되었다고 있었다고 있다고 있다고 있다고 있다면 보다 있다.		
	a) Antibiotic resistant ge	ne	b) Diseases resistant gene	e
	c) Allergy resistant gene		d) All of these	
8.		ucing plasmid (Ti) of <i>Agro</i>	bacterium tumefaciens is	used as a cloning vector.
	This statement is			
	a) True		b) False	
	c) Sometimes (a) and son		d) Neither (a) nor (b)	
9.		1. TO	e.g., ampicillin) is transferr	
		(1945) - 통일 - (1947)	e ampicillin resistant gene	
192920	a) Vectors	b) Plasmid	c) Selectable marker	d) Cloning sites
10.		ephalopathy disease is equa		
	a) Kala Azar		b) Parkinson's disease	
	c) Creutzfeldt-Jacob dise		d) None of the above	
11.	(95)	that is used to find comple		N = 1
	a) Vector	b) Plasmid	c) DNA probe	d) Recombinant DNA
12.	Proteins are removed by		X 0 W 1	D.B.
40	a) Ribonuclease	b) Chitinase	c) Cellulase	d) Protease
13.	- In Co.	AT(1),	he vector in genetic engine	Acres (Cress)
	a) It is resistant to antibi		b) It is resistant to restric	-
	c) Its ability to carry a for	MATERIAL STATE OF THE STATE OF	d) Its ability to cause infe	ction in the host
14.		py numbers of the linked D		l
			cycline resistance gene in t	
		3	ce due to insertion of foreig	אוט ח <u>ו</u>
	Choose regarding the abo		a) Dark and to	J) D-th C-1
1 =	a) I is true, II is false	b) II is true, I is false	c) Both are true	d) Both are false
15.		animai superior by view of	f genotype, introducing son	ie foreign genes in it, the
	phenomenon is called:			





- a) Tissue culture
- b) Biotechnology
- c) Genetic engineering
- d) Immunisation
- 16. Many copies of a DNA molecule in a test tube are produced by:
 - a) Polymerase chain reaction (PCR)
- b) Molecular chain reaction (MCR)
- c) Ephemeral chain reaction (ECR)
- d) All of them
- 17. Producing a 'giant mouse' in the laboratory was possible through: a) Gene mutation
 - b) Gene duplication
- c) Gene synthesis
- d) Gene manipulation

- 18. Downstream process includes
 - I. Separation of the product from the reactor
 - II. Purification of the product
 - III. Formation of the product with suitable preservatives
 - IV. Quality control testing and clinical trials in case of drugs

Which of the statements given above are correct?

- b) I, II and IV
- c) II, III and IV
- d) I, II, III and IV

- 19. More advancement in genetic engineering is due to
 - a) Restriction endonuclease

b) Reverse transcription

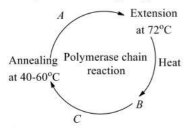
c) Protease

- d) Zymase
- 20. Plasmid are suitable vectors for gene cloning because
 - a) These are small circular DNA molecules, which can integrate with host chromosomal DNA
 - b) These are small circular DNA molecules with their own replication origin site
 - c) These can shuttle between prokaryotic and eukaryotic cells
 - d) These often carry antibiotic resistance genes
- 21. Polymerase chain reaction is useful in
 - a) DNA synthesis

b) DNA amplification

c) Protein synthesis

- d) Amino acid synthesis
- 22. Study the following diagram and identify A, B and C



- a) A-Tag polymerase, B-Denaturation at 94°C, C-Primer
- b) A-Denaturation at 94°C, B-Taq polymerase, C-Primer
- c) A-Primer, B-Denaturation at 94°C, C-Taq polymerase
- d) A-Taq polymerase, B-Extension, C-Transformation
- 23. A bioreactor is
 - a) Hybridoma

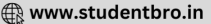
- b) Culture containing radioactive isotopes
- c) Culture for synthesis of new chemicals
- d) Fermentation tank
- 24. Which of the following techniques can be used to detect genetic disorders in human?
 - a) Polymerase Chain Reaction (PCR)
- b) Gel electrophoresis

c) Spectroscopy

- d) All of the above
- 25. Special sequence in the DNA recognized by restriction endonuclease is called
 - a) Restriction nucleotide sequence
- b) Palindromic nucleotide sequence
- c) Recognition nucleotide sequence
- d) All of the above

- 26. Primers are
 - a) Small chemically synthesized oligonucleotides of about 10-18 nucleotides that are complementary to the region of template DNA
 - b) Chemically synthesized oligonucleotides of about 10-18 nucleotides that are not complementary to the region of template DNA





	c) The double-stranded DNA that need to the ampl		
	d) Specific sequences present on recombinant DNA	A	
27.	This method of finding a gene is used when resea	rchers very little about the	gene they are trying to find.
	This process results in a complete gene library : a	collection of copies of DNA	fragments that represent the
	entire genome of an organism. Identify the method	l	
	a) Cloning b) Shotgun cloning	c) Gene synthesis	d) Cloning
28.	Consider the following statement about PCR		
	I. Polymerase Chain Reaction (PCR) is a technique	of synthesizing multiple co	pies of the desired gene in
	vitro		
	II. This technique was developed by Kary Mullis in	1985	
	III. A single PCR amplification cycle involves three	basic steps; denaturation, a	nnealing and extension
	Which of the statement given above are correct?		
	a) I and II b) I and III	c) II and III	d) I, II and III
29.	A somatic plant cell has potential to develop into a	full plant. This is called:	
	a) Totipotency b) Gene cloning	c) Tissue culture	d) Regeneration
30.	Ori is a DNA sequence that is responsible for initia	ting replication. This stater	nent is
	a) True	b) False	
	c) Sometimes (a) and sometimes (b)	d) Neither (a) nor (b)	
31.	Plasmids are autonomously replicating circular ext	trachromosomal DNA. This	statement is
	a) True	b) False	
	c) Sometimes (a) and sometimes (b)	d) Neither (a) nor (b)	
32.	Genetic engineering is possible because:		
	a) The phenomenon of transduction in bacteria is	well understood	
	b) We can see DNA by electron microscope		
	c) We can cut DNA at specific sites by endonucleas	es like DNA ase I	
	d) Restriction endonucleases purified form bacteri	a can be used in vitro	
33.	A single PCR amplification cycle involves		
	a) Denaturation b) Annealing	c) Extension	d) All of these
34.	DNA fingerprinting is related to:		
	a) Molecular analysis of profiles of DNA samples		
	b) Analysis of DNA samples using imprinting device		
	c) Techniques used for molecular analysis of differ		
	d) Techniques used in identification of fingerprints	s of different persons	
35.	The basic of DNA fingerprinting is:		
	a) The double helix	b) Errors in base sequer	ice
	c) Polymorphism in sequence	d) DNA replication	
36.	In genetic engineering, the terms vector is applied		201000
	a) Plasmid b) Sources of DNA	c) Cell which receives	d) Virus
37.	Which of the following are used to gene cloning?	12 12 12 12 12 12 12 12 12 12 12 12 12 1	
	a) Nucleoids b) Chromosomes	c) Mesosomes	d) Plasmid
38.	The process that preserves the distribution of DNA	a fragments in the gel while	creating replica on the filter
	is one of the following		
	a) Directed sequencing of BAC counting	b) Random shotgun seq	uencing
00	c) Electrophoresis	d) Southern blotting	
39.	Two enzymes responsible for restricting the growt		
	methylase and other was restriction endonuclease		
	a) Protection of host DNA from the action of restric		ng methyl group to one or
	two bases usually with in the sequence recogniz		
	b) Able to ligate the two cohesive ends of DNA mol		and a language
	c) Able to remove the methyl group and hence, pre	event the action of restriction	on endonuclease on host DNA

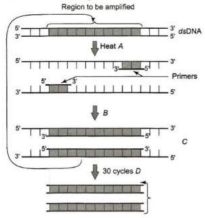
CLICK HERE >>

- d) Able to cut the DNA of bacteriophage at specific sites
- 40. Single-stranded DNA molecules that can bind to and be used to detect other DNA molecules are called
 - a) Primer
- b) STRs
- c) RFLPs
- d) Probes
- 41. Which of the following enzyme is used in genetic engineering?
 - a) Translocase

b) Topoisomerase

c) DNAse

- d) Restriction endonuclease
- 42. The below diagram refer to PCR. Identify the steps A, B and C and select the correct option



- A-Denaturation of 94-96°C, B-Annealing of 40-60°C, C-Extension through taq polymerase at 72°C, D-Amplified
- A-Annealing of 94-96°C, B-Denaturation of 40-60°C, C-Extension through *taq* polymerase at 72°C, D-Amplified
- A-Extension through taq polymerase at 40-60°C, B-Amplified, C-Denaturation of 40-60°C, D-Annealing of 94-96°C
- A-Annealing, B-Extension through taq polymerase at 40-60°C, C-Denaturation of 94-96°C, D-Annealing of 40-60°C
- 43. The controlled use of biological agents, such as microorganism, plants or animal cell, for beneficial use is called
- a) Biochemistry
- b) Molecular biology
- c) Biotechnology
- d) Microbiology

- 44. Humulin is a:
 - a) Pig insulin
- b) Human insulin
- c) Viral insulin
- d) Human clone

- 45. Find the incorrect statement:
 - a) Gene therapy is a genetic engineering technique used to treat disease at molecular level by replacing defective genes with normal genes
 - b) Calcitonin is a medically useful recombinant product in the treatment of infertility
 - c) Bt toxin is a biodegradable insecticide obtained from Bacillus thuringiensis
 - d) Trichoderma sp. is a biocontrol agent for fungal diseases of plants
- 46. Plasmids are extrachromosomal circular DNA molecules:
 - a) Which have their own point of replication and can replicate independently
 - b) Which have their own point of replication but cannot replicate independently
 - c) Which do not have their own point of replication and cannot replicate independent of bacterial of bacterial chromosomal DNA
 - d) None of the above
- 47. The genome map was produced under human genome project in:
 - a) 1992
- b) 1994
- c) 1996
- d) 2000

- 48. Term hybridoma implies:
 - a) DNA-RNA hybrid

b) Recombination of DNA molecules

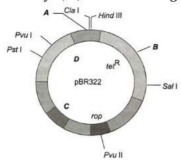
c) Somatic hybridisation

- d) Genetic fusion
- 49. Which of the following is a difficulty in getting prokaryotic cells to express eukaryotic genes?





- a) The signals that control gene expression are different and prokaryotic promoter regions must be added to the vector
- b) The genetic code differs between the two because prokaryotes substitute the base uracil for thymine
- c) Prokaryotic cells cannot transcribe introns because their genes do not have them
- d) The ribosomes of prokaryotes are not large enough to handle long eukaryotic genes
- 50. In transgenics, the expression of transgene in the target tissue is known by:
 - a) Enhancer
- b) Transgene
- c) Promoter
- d) Reporter
- 51. Identify A, B, C and D in the given diagram of E. coli cloning vector pBR322



- a) A-Eco RI, B-Bam HI, C-Ori, D-ampR
- b) A- ampR, B- Ori, C-Bam HI, D-Eco RI
- c) A-Ori, B-Bam HI, C-Eco RI, D-ampR
- d) A-Bam HI, B-Eco RI, C-ampR, D-Ori
- 52. Consider the following statements
 - I. In microinjection method foreign DNA is directly injected into the nucleus of animal cell or plant cell by using micro needles or micro pipettes
 - II. Microinjection method is used in oocytes, eggs and embryo
 - III. Electroporation is the formation of temporary pores in the plasma membrane of host cell by using lysozyme or calcium chloride
 - IV. In chemical mediated gene transfer method certain chemicals such as CO_2 help foreign DNA to enter the host cell

Which of the statements given above are correct?

- a) I and II
- b) I, II and III
- c) II, III and IV
- d) I, II, III and IV
- 53. The construction of the first recombinant DNA was done by using the native plasmid of:
 - a) E. coli

b) Salmonella typhimuriumd) Yeast

- c) B. thuringiensis
 - a) Microinjection

b) ELISA

c) Polymerase chain reaction

- d) Gene gun
- 55. Polyethylene glycol method is used for
 - a) Biodiesel production

- b) Seedless fruit production
- c) Energy production from sewage
- d) Gene transfer without a vector
- 56. The enzymes, commonly used in genetic engineering are
 - a) Restriction endonuclease and polymerase

54. Gene amplification using primers can be done by

- b) Endonuclease and ligase
- c) Restriction endonuclease and ligase
- d) Ligase and polymerase
- 57. Which one of the following techniques had helped to solve many mysteries involving murders, robberies and rapes?
 - a) Gene splicing

b) Computer technology

c) DNA fingerprinting

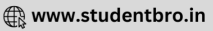
- d) Gene cloning
- 58. Consider the following statements
 - I. Recombinant DNA technology popularly known as genetic engineering is a stream of biotechnology which deals with the manipulation of genetic material by man *in vitro*
 - II. pBR322 is the first artificial cloning vector developed in 1977 by Boliver and Rodriquez from $E.\ coli$ plasmid





	III. Restriction enzymes belongs to a class of enzymes called nucleases				
	Which of the statements	given above are correct?			
	a) I and II	b) I and III	c) II and III	d) I, II and III	
59.	What is C-DNA?				
	a) Circular DNA				
	b) Cloned DNA				
	c) DNA produced from re	everse transcription of RNA	\		
	d) Cytoplasmic DNA				
60.	PCR was developed by	A inB and for this he	received Nobel Prize for cl	nemistry inC Here A, B	
	and C can be recognized				
	A B C				
	a) Kary Mulllis 1990 19	997	b) Flemming 1985 1	993	
	c) Kary Mullis 1985 1		d) Flemming 1990		
61.		from a plasmid was done v	,		
	B Here A and B can b			, , ,	
	a) A-Tu ligases; B-Molecu		b) A-Restriction enzyme	: B-Molecular scissors	
	c) A-Joining enzyme; B-M	•	d) A-DNA polymerases;		
62.	, , , , , , , , , , , , , , , , , , , ,	experiment, restriction enz			
25.76	a) Bacterial DNA only		b) Viral DNA only		
	c) Any DNA fragment		d) Eukaryotic DNA only		
63.	The components of a bio	reactor are	-,,		
	I. an agitator system				
	II. an oxygen delivery sys	tem			
	III. foam control system				
	IV. temperature control s	vstem			
	V. pH control system	y			
	The same of the sa	h draw cultures periodicall	v		
	Choose the correct option	270	*		
	a) I, II, III, IV and V	b) II, IV, V and VI	c) I, II, III, IV and VI	d) All of these	
64.		istron in base pairs which:		of 50 amino acids is:	
	a) 50 bp	b) 100 bp	c) 150 bp	d) 200 bp	
65.	I. DNA being a hydrophili	ic molecule cannot pass thr	ough cell membranes	*	
	II. The bacteria should be	made competent to accept	t the DNA molecule		
	The correct option regard	ding the above statements	is		
	a) I is true, but II is false		b) II is true, but I is false		
	c) I and II are true		d) I and III are false		
66.	In cloning plasmid pBR32	22			
	p stands forA				
	B stands forB				
	R stands forC				
	Choose the correct option	n			
	a) A-plasmid, B-Boliver, (C-Rodriquez	b) A-plasmid, B-bacteria	, C-Rodriquez	
	c) A-prophage, B-bacteri	ophage, C-Rodriquez	d) A-prophage, B-Bolive	r, C-Rodriquez	
67.	Blood stains are found a	t the site of murder. If DN	A profiling technique is t	o be used for identifying the	
	criminal, which of the fol	lowing is ideal for use?			
	a) Serum	b) Erythrocytes	c) Leucocytes	d) Platelets	
68.	The Ti plasmid used in ge	enetic engineering is obtain	ned from:	<i>8</i> 0	
	a) Bacillus thuringeinsi	S	b) Agrobacterium rhizo	ogenes	
	c) Agrobacterium tumij	^f aciens	d) Escherichia coli		





69.	Who got the Nobel prize	in medicine for their disco	very of 'G-proteins' and the	role of these proteins in the
	cells:			.
	a) Robert and Philip Shar	ъ	b) Gilman and Rodbell	
	c) Fischer and Krebs		d) Ervin Nahar and Bert	Sakmann
70.	Which of the following is	required to perform polyn	nerase chain reaction?	
	I. DNA template			
	II. Primer			
	III. Taq polymerase and 1	77 (770)		
	Choose the correct option			
	a) I, II and III	b) I and II	c) II and III	d) II and III
71.	The basis for DNA finger			
		on fragment length polym	orphism (RFLP)	
	b) Phenotypic differences			
	c) Availability of cloned I			
70	d) Knowledge of human l	5 5 5		1
72.		의 Banka Bellem (1975년 1981년) 원인 원인 원인 경험 경영 (1984년) 원인 1984년 1987년	erest, is transferred to the	- 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
	more of these can be used		ard and select the correct o	ption about which one or
	I. Bacterium	u as a vector/vectors		
	II. Plasmid			
	III. Plasmodium			
	IV. Bacteriophage			
	a) I, II and IV	b) I only	c) I and III	d) II and IV
73.			anism can be done through	450
	a) Genetic engineering	b) Tissue culture	c) Transformation	d) None of these
74.			m thermophilic bacteria ar	
	a) DNA polymerase		b) DNA ligases	
	c) Restriction endonucle	ases	d) RNA polymerases	
75.	Due to chloramphenicol i	resistance gene, one is able	to select a transformed cel	ll in the presence of
	THE STATE OF THE S	oramphenicol resistance g		
	a) Origin of replication		b) Selectable marker	
	c) Cloning sites		d) Insertional inactivatio	n
76.	100 - 100 -	n site for the restriction en		200000000000000000000000000000000000000
	a) <i>Eco</i> RI	b) <i>Hind</i> II	c) <i>Eco</i> RII	d) <i>Bam</i> HI
77.	Plasmid is	:		
		icating circular extrachron		
	c) An circular protein mo	icating circular extrachron	1050IIIai KIVA	
		icating chromosomal DNA		
78		action is a technique that is		
, 0.	a) <i>In vivo</i> replication of I	1.9	, asca for	
	b) <i>In vivo</i> synthesis of <i>m</i> l			
	c) <i>In vitro</i> synthesis of <i>m</i>			
			ng thermostable DNA polyn	nerase
79.		ant in genetic engineering		
	a) Has plasmids that can	be genetically engineered		
	b) Allows the study of eu	karyotic gene regulation a	nd expression	
	c) Grows readily and rap	idly in the laboratory		
	d) All of the above			
80.	The genome of Caenorho	abditis elgans consists of:		

	a) 3 billion base pairs and 30,000 genes	b) 12 million base pairs a		
	c) 4.7 million base pairs and 4,000 genes	d) 97 million base pairs a	nd 18,000 genes	
81.				
	a) Nitrosomonas and Klebsiella	b) Escherichia and Agro	bacterium	
	c) Nitrobacter and Azotobacter	d) Rhizobium and Diploc	roccus	
82.	Gel electrophoresis is used for:			
	a) Isolation of DNA molecule			
	b) Cutting of DNA into fragments			
	c) Separation of DNA fragments according to their s	size		
	d) Construction of recombinant DNA by joining with	n cloning vectors		
83.	Then linking of antibiotic resistance gene with the p	olasmid vector became poss	ible with:	
	a) DNA ligase b) Exonucleases	c) Endonucleases	d) DNA polymerase	
84.	Restriction endonucleases are:			
	a) Present in mammalian cell for degradation of DN	A when the cell dies		
	b) Synthesized by bacteria as part of their defence r	nechanism		
	c) Used for in vitro DNA synthesis			
	d) Both (B) and (C)			
85.	Which one of the following is related with genetic e	ngineering?		
	a) Plasmids b) Mitochondria	c) Mutations	d) Ribosomes	
86.	Enzyme that is used in PCR technology is			
	a) Ligase	b) Polymerase		
	c) Helicase	d) Reverse transcriptase		
87.	Genetic diagnosis by DNA testing:			
	a) Detects only mutant and normal alleles			
	b) Can be done only on eggs or sperms			
	c) Involves hybridization to ribosomal RNA			
	d) Utilizes restriction enzymes and a polymorphic s	ite		
88.	An enzyme catalyzing the removal of nucleotides from	om the ends of DNA is		
	a) Endonuclease b) Exonuclease	c) DNA ligase	d) <i>Hind</i> II	
89.	Inducible/lac operon system operates in:			
	a) Catabolic pathway	b) Anabolic pathway		
	c) Intermediate metabolism	d) All the above		
90.	Polymerase Chain Reaction (PCR) needs			
	a) DNA template b) Primers	c) Taq polymerase	d) All of these	
91.	Consider the following statements			
	I. A soil inhabiting plant bacterium, Agrobacterium	tumefaciens, a pathogen	of several dicot plants is	
	able to transfer a piece of DNA known as T-DNA			
	II. The T-DNA causes tumours			
	III. Tumour formation induced by Ti-plasmid			
	Which of the statements given above are correct?			
	a) I and II b) I and III	c) II and III	d) I, II and III	
92.	Restriction endonucleases are enzymes which			
	a) Make cuts at specific positions within the DNA m			
	b) Recognize a specific nucleotide sequence for bind			
	c) Restrict the action of the enzyme DNA polymeras			
	d) Remove nucleotides from the ends of the DNA m	olecule		
93.	Restriction enzymes are used to cut			
	a) Single-stranded RNA	b) Double-stranded DNA		
	c) Single-stranded DNA	d) Double-stranded RNA		
94.	Restriction enzymes are isolated chiefly from:			

			2.5	
0.5	a) Algae	b) Fungi	c) Protozoans	d) Prokaryotes
95.	Which of the following is		15.77	D.
	a) Agrobacterium tumefa	iciens – Tumour	b) Thermus aquaticus – I	
0.0	c) pBR322 - Enzyme	(PCP)	d) Ligase – Molecular sci	
96.			in which amplification of sp	pecific DNA sequences is
	carried out in vitro. This	statement is	12.5.1	
	a) True	(1.)	b) False	
07	c) Sometimes (a) and sor	1977 N	d) Neither (a) nor (b)	
97.			nome is estimated to be ab	
00	a) 35 million	b) 3.1 billion	c) 3.5 million	d) 35 thousand
98.	Totipotency in cell is:	d:,,,,,		
	a) Flower in a culture me		adium	
		rom a flower in a culture m ganism from cell in culture		
		sues of all kinds from a cell		
99	Restriction enzymes was		in a culture medium	
,,,	a) Alexander Flemming	discovered by	b) Waksman	
	c) Berg		d) Smith, Nathan and Ark	ner
100	. Identify the plasmid:		aj Simini, Natilan ana mi	, c.
100	a) Alu I	b) Hind III	c) Eco RI	d) pBR 322
101	. Consider the following st		c) 200 m	a) p2.1022
			n raw materials are biologi	cally converted into specific
	products			
	•	only used bioreactors is of	stirring type	
		ar paramet e n na maiori da espera da la composita de la composita de la composita de la composita de la composita Mos	e desired materials on a sm	all scale in the laboratory
		7.75 U.75 1275	ical product is done by usir	7/
	a) I and II	b) I and III	c) I, II and III	d) I, II, III and IV
102	. The term 'Biotechnology'	was given by	2724 93	67% 59 (800)
	a) Craig Venter	b) Robert Edward	c) Karl Erkey	d) Temin and Baltimore
103	. A collection of organisms	, usually viruses, bacteria o	or yeast, which have been t	ransformed by the addition
	of extra genes from anoth			
	 a) Gene replication 	b) Gene cloning	c) Gene pool	d) Gene library
104	_		m the end of the polynucled	otide chain are:
	a) Specific for 5' end of R			
	b) Specific for 3' end of R			
		l 3' ends of nucleotide strai		
01202		3' ends of nucleotide strar		07 (324) N
105			plasmid into 1 phage genon	
400	a) Cosmid	b) Phasmid	c) Phagmid	d) Foreign DNA
106	. Which of the following st		.1 . 1 . 1 1	. 1 6 11 11
			NATURE - ANALYS IN <mark>T</mark> ERNATION INSIDE WITH INTERNATION OF SALE	vas taken from an udder cell
		s appeared first in dinosau	rs	
	c) Heart of mammals is in	and a file of the court of the	v.	
107		upright in pond ecosystem	l.	
107	. Which of the following st		rough call mambranes	
	NS(15) S	c molecule cannot pass thr	DNA known as 'Z-DNA' in t	he Ti-nlasmid which
	no-transcorperities are concerned to transcorperities and the confidence are confidence and the confidence are con-		roduce chemical against pa	angger at the same section of the was a surface to the same section of the same sectio
	(5)	58:		s in animal because of their
		al cells into cancerous cell		5 m ammar because of thell
	ability to transform norm	ar cens into cancerous tell		

	eering, DNA from different sour		e restriction enzymes so that
~	s have same kind of sticky ends	i e	
Choose the correct	option		
a) Only I	b) Only II	c) Only III	d) Only IV
108. Which one of the fo	llowing pairs is correctly match	ied?	
a) RNA polymerase	-RNA primer	b) Restriction enzyme	s-Genetic Engineering
c) Central Dogma-c	odon	d) Okazaki fragments-	
109. Bam HI, Eco RI, Sma			• • • • • • • • • • • • • • • • • • • •
a) Restriction endo	1,200	b) Restriction endonu	cleases
c) Restriction exon		d) Restriction polyme	
110. PCR technique was		-,	
a) Boyer	b) Kary Mullis	c) Cohen	d) Sanger
111. Somaclonal variation	Control of the contro	e, donen	a) banger
a) Hybridization	in can be obtained by.	b) Tissue culture	
c) Application of co	lahiaina	d) Irradiation with gai	mma vave
3 1517		u) ii radiation with gai	illilla rays
112. Ability to absorb for		a) 1164	d) Transadoration
a) Sexduction	b) Competence	c) Hfr	d) Transduction
	ing is specifically used in genet		
a) Ligase		b) Gyrase	
c) DNA polymerase		d) Restriction endonu	
	ing capacity of Agrobacteriu	m tumefaciens is locate	ed in large extrachromosomal
plasmids called			
a) Ri-plasmid	b) Lambda phage	c) pBR322	d) Ti-plasmid
115. Who discovered red	combinant DNA (rDNA) techno	logy?	
a) Har Gobind Khor	ana	b) James D Watson	
c) Stanley Cohen an	d Herber Boyer	d) Walter Sutton and	Avery
116. Which of the follow	ing is used in recombinant DNA	A technique?	
a) Cell wall of virus		b) Gene which produc	es capsid of virus
c) Virus		d) Capsid of virus	
117. There are special pr	oteins that help to open up DN	A double helix infront of	the replication fork. These
proteins are:			
a) DNA gyrase	b) DNA polymerase I	c) DNA ligase	d) DNA topoisomerase
	rom sea weeds finds use in:	-)	,
a) Spectrophotome		b) Tissue culture	
c) Gel electrophore		d) PCR	
119. For selectable mark		u) i ok	
	he host cells which contain the	vector and eliminate the	non transformants
			, tetracycline or kanamycin, are
useful selectable ma		picinin, cinoramphenicor	, tetracycline or kanamychi, are
	nents given above are correct?	-> 1 1 11	D.N Cal
a) Only I	b) Only II	c) I and II	d) None of these
120. The first clone anim			
a) Molly sheep	b) Polly sheep	c) Dolly sheep	d) Molly goat
	used in genetic engineering is:		
a) E.coli	b) Diplococcus	c) Rhizobium	d) Spirillium
	it restriction enzymes have the		strands in a particular
	vhat has became known as 'stic		
a) Ramdeo Mishra	b) Stanley Cohen	c) Herbert Boyer	d) James D Watson
123. A restriction fragme	ent containing a specific gene o	f interest can be identifie	d by gel electrophoresis
followed by transfe	rring the DNA to a membrane a	s a solid support matrix ι	ısing a procedure called



a) An allozyme

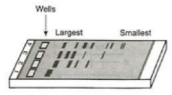
b) A southern blot

c) Identification of a gene

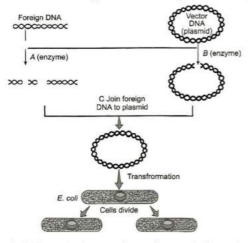
d) A restriction fragment length polymorphism

- 124. About gene gun method
 - I. This method is also known as biolistic technique
 - II. In this method cells are bombarded with high velocity micro-particles of gold or tungsten coated with DNA in plants
 - III. Important crop plants like maize, rice and wheat have now been transformed by this method Which of the statements given above are correct?
 - a) I and II
- b) I and III
- c) II and III
- d) I, II and III

125. Identify the correct match for the given diagram



- a) Electrophoresis Migration of undigested and digested set of DNA fragments
- b) Bioreactor Raw materials are biologically converted into specific products
- c) Microinjection Technique of introducing foreign genes into a host cell
- d) Gene cloning Technique of obtaining identical copies of a particular DNA segment
- 126. In DNA fingerprinting which of the following is true?
 - a) VNTR is used as probes
 - b) Specific metabolic genes are used as probes
 - c) House keeping or luxury genes are use as probes
 - d) All of the above
- 127. The message from nuclear DNA for the synthesis of specific cytoplasmic protein is carried by:
 - a) mRNA
- b) t-RNA
- c) s-RNA
- d) r-RNA
- 128. The recent techniques used for separating fragments of DNA is:
 - a) Northern blotting
- b) Southern blotting
- c) Eastern blotting
- d) Western blotting
- 129. The flowchart given below represent the process of recombinant technology. Identify A and D



- a) A-Restriction endonuclease, B-Restriction exonuclease, C-RNA ligase, D-Transformation
- b) A-Restriction endonuclease, B-Restriction endonuclease, C-DNA ligase, D-Transformation
- c) A-Restriction exonuclease, B-Restriction endonuclease, C-DNA polymerase, D-Transduction
- d) A-Restriction endonuclease, B-Restriction endonuclease, C-DNA polymerase, D-Transformation
- 130. RNA is removed by the treatment with
 - a) Ribonuclease
- b) Protease
- c) Chitinase
- d) Cellulase
- 131. Which one of the following scientists developed the process of DNA fingerprinting?
 - a) Kary B. Mullis
- b) T.H. Morgan
- c) H.O. Smith
- d) Alec Jeffreys



132.	132. Which of the following statement is not correct regarding <i>Eco</i> RI restriction endonuclease enzyme?				
102.	I. Eco. RI restriction endonuclease enzyme is isolated from Escherichia coli RY13				
	II. Its recognition sequence is 5'-GAATTC - 3'	ed if on Escher tenta con K	.13		
	3'-CTTAAG - 5'				
	3 -C11AAG - 3				
	, , , , , , , , , , , , , , , , , , ,				
	5' - G - A - A -T - T - C - 3'				
	III. Its site of cleavage is				
	3' - C - T - T - A - A - G - 5				
	1				
	a) I and II				
	b) I and III				
	c) I, II and III				
	d) None of the above				
133.	Process of formation of RNA from DNA is called				
	a) Transduction b) Transcription	c) Transformation	d) Translation		
134.	Which of the following would not be used in prepar	ring recombinant DNA?			
	a) Plasmids	b) Phages			
	c) Restriction enzymes	d) DNA polymerase III			
135.	Which one of the following bacteria has found exten	and the control of th	ring work in plants?		
	a) Agrobacterium tumefaciens	b) Clostridium septicum			
	c) Xanthomonas citri	d) Bacillus coagulens			
136	Which of the following components are used in gel	· ·			
100.	I. Ethidium bromide	ciccii opnoresis:			
	II. Restriction endonuclease				
	III. Agarose				
	IV. UV radiation				
	Choose the correct option	-) I II I IV	4) 1 11 111 4 117		
107	a) I and II b) I and III	c) I, II and IV	d) I, II, III and IV		
13/.	What is the first step in Southern Blotting techniqu				
	a) Isolation of DNA from a nucleated cell such as th		ne		
	b) Denaturation of DNA on the gel for hybridization				
	c) Production of group of genetically identical cells				
	d) Digestion of DNA by restriction enzyme	14 300 000 B00 4 57407			
138.	The most thoroughly studied of the known bacteria				
	a) Plant growth simulation by phosphate-solubilisi	1700 C			
	b) Cyanobacterial symbiosis with some aquatic ferr				
	c) Gall formation on certain angiosperms by Agrob				
	d) Nodulation of Sesbania stems by nitrogen fixing	bacteria			
139.	Microorganisms can be grown in the bioreactor by				
	a) Support growth system	b) Agitated growth system	m		
	c) Suspended growth system	d) Both (a) and (b)			
140.	In Northern blotting RNAs are separated by gel elec	ctrophoresis and the RNA ba	ands are transferred onto a		
	membrane of:				
	a) Diazobenzyl oxymethyl	b) Diazobenzene			
	c) Diazobromine	d) None of the above			
141.	Which one of the following is commonly used in tra		rop plants?		
	a) Trichoderma harzianum	b) <i>Meloidogyne incogniti</i>	_ 5 S		
	c) Agrobacterium tumefaciens	d) <i>Penicillium expansum</i>			
142	Which one among the following is just a cloning pla				
1578678770	a) pBAD-18-Cam b) pBCSK	c) pUC 18	d) pET		
		Sector Control			

143. ThereA are the DNA molecules that can carry a	foreignB segment into	the host cell.
Here A and B refers to		
A B	h) Waston DNA	
a) Vector RNA c) Gene RNA	b) Vector DNA d) Gene DNA	
See Presentative Constitution	d) Gene DNA	
144. Probes, used in DNA fingerprinting are initially	h) Mini catallita	
a) Single-stranded RNA	b) Mini satellite	
c) 19 base long oligonucleotides145. Application of PCR are	d) All of the above	
I. detection of pathogens		
II. diagnosis of specific mutation		
III. DNA fingerprinting		
Choose the correct option		
a) I and II b) I and III	c) II and III	d) I, II and III
146. A clone of sheep Dolly has been made by:	c) ii ana iii	a) i, ii ana iii
a) Gene transfer	b) Somatic cell cloning	
c) Nucleus transfer	d) Germinal cell cloning	
147. T ₁ -plasmid used in genetic engineering is obtained in		
a) Bacillus thuringiensis	b) <i>Agrobacterium rhizog</i>	renes
c) Agrobacterium tumefaciens	d) Psedomonas syringae	
148. The role of DNA ligase in the construction of a recor		
a) Formation of phosphodiester bond between two		
b) Formation of hydrogen bonds between sticky en		
c) Ligation of all purine and pyrimidine bases	200	
d) None of the above		
149. Transgenic animals are produced by injecting foreign	gn gene into the:	
a) Egg	b) Nucleus of unfertilized	d egg
c) Nucleus of fertilized egg	d) Nucleus of sperm	.55.57
150. Clonal cell lines can be obtained by:	Visit Productive Annual Control Contro	
a) Autoradiography b) Tissue culture	c) Centrifugation	d) Cell fractionation
151. Electroporation procedure involves:		
a) Fast passage of food through sieve pores in phloe	em elements with the help	of electric stimulation
b) Opening of stomatal pores during night by artific	ial light	
c) Making transient pores in the cell membrane to i	ntroduce gene constructs	
d) Purification of saline water with the help of a me	mbrane system	
152. Which of the following is associated with genetic en	gineering?	
a) Plastids b) Plasmids	c) Mutations	d) Hybrid vigour
153. Biolistics (gene gun) is suitable for		
a) Disarming pathogen vectors	b) Transformation of pla	nts cells
c) Construction recombinant DNA by joining with	d) DNA fingerprinting	
vectors		-2
154. Which of the following statements are correct for the	ie enzyme <i>taq</i> polymerases	S.f.
I. Taq polymerase is thermally unstable	Cu alama animati an	
II. It requires primers for carrying out the process o		
III. <i>Taq</i> polymerase is isolated from thermophilic ba	acterium, i nermus aquati	cus
Choose the correct option	a) II and III	d) I II and III
a) I and II b) I and III 155. EFB stands for	c) II and III	d) I, II and III
a) European Federation of Biotechnology	b) Eurasian Federation o	of Riotechnology
c) East Asia Federation of Biotechnology	d) Ethiopian Federation	
c) East Asia redefation of biotechnology	a) Eunopian rederation	or protectifiology

To or time commonly about Divitinger printing recommende	in forensic studies is simpl	y a method involving		
a) Southern blotting b) Northern blotting	c) Eastern blotting	d) Western blotting		
157. Cry I endotoxins obtained from Bacillus thruigien	usis are effective against			
a) Nematodes b) Bollworms	c) Mosquitoes	d) Flies		
158. In the naming of restriction enzymes the first letter	r is derived fromA nam	e and next two letters from		
theB and fourth letter fromC ofD wher	e the enzymes are extracted	1:		
A to D in the statement can be				
A B C D				
a) Genus species strain bacteria	b) Species genus strain	bacteria		
c) Genus species variety eukaryote	d) Species genus variet	y eukaryote		
159. Which of the following techniques is most common	lly used to separate DNA m	olecules by size?		
a) Chromatography b) PCR	c) RFLP	d) Gel electrophoresis		
160. Which one of the following scientists got the Nobel	Prize for his invention poly	merase chain reaction		
(PCR)?				
a) F. Sanger b) Paul Berg	c) Alec Jeffreys	d) Kary B. Mullis		
161. Which is non-invasive technique of genetic counse	TO STATE OF THE PARTY OF THE PA			
a) Amniocentesis	b) Chorionic biopsy			
c) Foetal blood sampling	d) Ultrasonography			
162. The colonies of recombinant bacteria appear white	in contrast to blue colonie	s of non-recombinant		
bacteria because of:				
 a) Insertional inactivation of alpha-galactosidase in 		í		
 b) Insertional inactivation of alpha-galactosidase in 				
c) Inactivation of glycosidase enzyme in recombina				
d) Non-recombinant bacteria containing beta-galac				
163. Which of the following steps are catalyzed by <i>taq</i> p	- 1989			
a) Denaturation of template DNA	b) Annealing of primers	to template DNA		
c) Extension of primer end on the template DNA				
164. I. In the process of recombinant DNA technology af	ter several treatment the p	urified DNA is precipitated		
by adding chilled ethanol	I DNA	1 'd DMA . '		
II. The bacterial/plant, animal cell is broken down by enzymes to release DNA, along with RNA, proteins,				
	by chaymes to release bring			
polysaccharides and lipids	by chaymes to release 51111			
polysaccharides and lipids Choose the correct option for above statements				
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false	b) I is false, but II is true			
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true	b) I is false, but II is true d) I and II are false			
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact	b) I is false, but II is true d) I and II are false ors?	w providing optimal growth		
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie	b) I is false, but II is true d) I and II are false ors? ving the desired product b	by providing optimal growth		
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit	b) I is false, but II is true d) I and II are false ors? ving the desired product b amin and oxygen			
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor	b) I is false, but II is true d) I and II are false ors? ving the desired product b amin and oxygen			
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor Correct option is	b) I is false, but II is true d) I and II are false fors? ving the desired product b amin and oxygen ganisms under aseptic cond	litions for a number of days		
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor Correct option is a) Only I b) Only II	b) I is false, but II is true d) I and II are false fors? ving the desired product b amin and oxygen ganisms under aseptic cond c) I and II			
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor Correct option is a) Only I b) Only II 166. Taq polymerase enzyme used in PCR is isolated from	b) I is false, but II is true d) I and II are false fors? ving the desired product b amin and oxygen ganisms under aseptic cond c) I and II	litions for a number of days d) None of the above		
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor Correct option is a) Only I b) Only II 166. Taq polymerase enzyme used in PCR is isolated from a) Thermus aquaticus	b) I is false, but II is true d) I and II are false fors? ving the desired product b amin and oxygen ganisms under aseptic cond c) I and II om b) <i>Thermococcus litoral</i>	litions for a number of days d) None of the above		
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor Correct option is a) Only I b) Only II 166. Taq polymerase enzyme used in PCR is isolated from a) Thermus aquaticus c) Salmonelia typhimurium	b) I is false, but II is true d) I and II are false fors? ving the desired product be amin and oxygen ganisms under aseptic cond c) I and II om b) Thermococcus litoral d) None of the above	litions for a number of days d) None of the above		
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor Correct option is a) Only I b) Only II 166. Taq polymerase enzyme used in PCR is isolated from a) Thermus aquaticus c) Salmonelia typhimurium 167. The first hormone artificially produced by culturing	b) I is false, but II is true d) I and II are false fors? ving the desired product be amin and oxygen ganisms under aseptic cond c) I and II om b) Thermococcus litoral d) None of the above g bacteria is:	litions for a number of days d) None of the above is		
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor Correct option is a) Only I b) Only II 166. Taq polymerase enzyme used in PCR is isolated from a) Thermus aquaticus c) Salmonelia typhimurium 167. The first hormone artificially produced by culturing a) Insulin b) Thyroxine	b) I is false, but II is true d) I and II are false fors? ving the desired product be amin and oxygen ganisms under aseptic cond c) I and II om b) Thermococcus litoral d) None of the above	litions for a number of days d) None of the above		
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor Correct option is a) Only I b) Only II 166. Taq polymerase enzyme used in PCR is isolated from a) Thermus aquaticus c) Salmonelia typhimurium 167. The first hormone artificially produced by culturing a) Insulin b) Thyroxine	b) I is false, but II is true d) I and II are false fors? ving the desired product be amin and oxygen ganisms under aseptic cond c) I and II om b) Thermococcus litoral d) None of the above g bacteria is: c) Testosterone	d) None of the above is d) Adrenaline		
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor Correct option is a) Only I b) Only II 166. Taq polymerase enzyme used in PCR is isolated from a) Thermus aquaticus c) Salmonelia typhimurium 167. The first hormone artificially produced by culturing a) Insulin b) Thyroxine 168. A gene is made up of: a) DNA b) RNA	b) I is false, but II is true d) I and II are false fors? ving the desired product be amin and oxygen ganisms under aseptic cond c) I and II om b) Thermococcus litoral d) None of the above g bacteria is: c) Testosterone c) Either DNA or RNA	d) None of the above is d) Adrenaline d) Amino acids		
polysaccharides and lipids Choose the correct option for above statements a) I is true, but II is false c) I and II are true 165. Which of the statements are correct about bioreact I. It provides all the optimal conditions for achie conditions like temperature, pH, substrate, salt, vit II. It is suited for large-scale production of microor Correct option is a) Only I b) Only II 166. Taq polymerase enzyme used in PCR is isolated from a) Thermus aquaticus c) Salmonelia typhimurium 167. The first hormone artificially produced by culturing a) Insulin b) Thyroxine	b) I is false, but II is true d) I and II are false fors? ving the desired product be amin and oxygen ganisms under aseptic cond c) I and II b) Thermococcus litoral d) None of the above g bacteria is: c) Testosterone c) Either DNA or RNA s isolated by Smith, Wilcox	d) None of the above is d) Adrenaline d) Amino acids and Kelley fromB		

CLICK HERE >>

A B C a) Eco RI Escherichia RY 13 Restriction sequence b) Eco RII E. coli R 245 Recognition sequence c) Hind II Haemophilus influenza Recognition sequence d) Bam HI Bacillus Restriction sequence amyoliquefaciens 170. In gel electrophoresis, the separated DNA fragments are visualized after staining the DNA with ... A... followed by exposure to ...B... Here A and B refers to Infrared radiation a) B-galactosidase b) Ethidium bromide UV radiation c) Ethidium nitrate d) Ethidium chloride Radiowave y-rays 171. In DNA fingerprinting: a) A positive identification can be made b) Multiple restriction enzyme digests/generate unique fragments c) The polymerase chain reaction amplifies fewer DNA d) The variability of repeated sequences between two restriction sites is evaluated 172. Cosmid is: a) Extragenetic material in Mycoplasma b) Circular DNA in bacteria c) Extra DNA in bacteria d) Fragment of DNA inserted in bacteria for forming 173. Following enzymes/chemical/technique are used in the process of gel electrophoresis I. sample DNA is cut into fragments II. restriction endonucleases III. agarose gel IV. ethidium bromide V. UV-radiation VI. elution Mark the correct sequence of their use a) I, II, III, VI, V and IV b) I, II, III, VI, V and IV c) IV, V, VI, I, II and III d) I, II, IV, V, VI and III 174. Improvement of genotype of an organism by addition of some foreign genes is: a) Genetic diversity b) Gene handling c) Tissue culture d) Genetic engineering 175. Which one is a true statement regarding DNA polymerase used in polymerase chain reaction? a) DNA polymerase is responsible for DNA synthesis b) It is isolated from Protozoa c) It is serves as a selectable marker d) It is used to ligate introduced DNA in recipient plant cell 176. Most sensitive technique to detect malignant cell in non-hodgkin's lymphoma is b) Gene therapy a) Polymerase chain reaction c) Stem cell therapy d) None of the above 177. Gene therapy involves: a) Introducing of a normal genes in cell b) Eliminating defective and useless genes c) Treating of defective genes with radiations d) Replacement of defective genes by normal ones 178. Human Genome project was the thought of: a) Jean Dausset b) Watson c) Crick d) None of the above 179. Which conserved motifs are found in *E. coli* genes? a) TATA box b) CAAT box c) Pribnow box d) All of these

100 Ci h-li	- CDNA -t d -ii th - h	
180. Given below is a sample of a portion is so special shown in it?	of DNA strand giving the base so	equence on the opposite strands. What
5'GAATTC3'		
3'S'		
a) Replication completed	b) Deletion mu	tation
c) Start codon at the 5' end		sequence of base pairs
181. The DNA used as a carrier for transfe		(A)
a) Cloning vector b) Vehicle		
182. The nuclease enzyme, which beings i	15%	
a) Exonuclease b) Kinase	c) Polymerase	
183. Genetically engineered bacterium use	ed in production of:	
a) Thyroxine b) Human	insulin c) Epinephrine	d) Cortisol
184. In Southern blotting is separate	d by gel electrophoresis:	
a) DNA b) m-RNA	c) t-RNA	d) Protein
185. Taq polymerase enzyme is found in:		
a) Thermus aquatecus b) E. coli	c) Pseudomon	A NAME OF THE PARTY OF THE PART
186. The process used for separation of pr		
	rn blotting c) Western blo	
187. Which of the following methods(s) is		
	trophoresis c) Elution	d) Extension
188. The figure shown three steps (A, B, C)		n PCR. Select the option giving correct
identification together with what rep	resents?	
A 3'	3' dsDNA	
5'	3'	
B 5' 5'	A	
3'	5'	
5'	3'	
C 3'		
3'	5′	
a) B-denaturation at a temperature of	of about 98°C separating the two	DNA strands
b) A-denaturation at a temperature of	f about 50°C	
c) C-extension in the presence of hea	t stable DNA polymerase	
d) A-annealing with three sets of prir	mers	
189. DNA fingerprinting method is very us		
 a) DNA tests for identity and relation 	17.	
c) Polymorphism	d) All of the ab	ove
190. Restriction endonucleases are used a	S:	
a) Molecular build up at nucleotides	3	
b) Molecular degradation to DNA bre		
c) Molecular knives for cutting DNA	State of the state	
d) Molecular cement to combine DNA		dust undergoes Consention and
191. After completion of the biosynthetic purification processes, collectively te		duct undergoes. Separation and
a) Transformation	b) Transductio	n
c) Downstream processing	d) Upstream p	
192. Which of the following should be cho		(477)
enzyme on a large scale, using microl		produce a recombinant protein or
a) Stirred-tank bioreactor	b) Electrophor	esis
c) Laboratory flask of largest capacit		
z, zazorator, mani or mi Boot cupacit	, aj im oi die do	

193. Go through the figure and select the option for C and D. Here A and B are taken as vector/plasmid DNA and foreign DNA respectively



Sticky end			
Restriction enzyme	Enzyme joining the		
recognizing palindrome (
a) <i>Eco</i> RI	DNA ligase	b) DNA ligase	<i>Eco</i> RI
c) Exonuclease	DNA ligase	d) DNA ligase	Exonuclease
194. Which of the following is			
a) Ligase		b) Polymerases	
c) Restriction endonucle	ases	d) Transcriptase	
195. A kind of biotechnology i		of DNA is	
a) DNA replication	b) Genetic engineering	g c) Denaturation	d) Renaturation
196. Harris and J.F. Watkins ir	n 1965 first time reporte	ed the fusion of following cell	lines to form hybrids:
a) Mouse and man		b) Mouse and hamster	
c) Mouse and click eryth	rocytes	d) Mouse and Drosophi	la
197. Polymerase chain reaction	on employs		
 a) Primers and DNA ligas 	se	b) DNA ligase only	
c) DNA polymerase		d) Primer and DNA poly	merase
198. An antibiotic resistance g	gene in a vector usually l	nelps in the selection of	
a) Competent cells	b) Transformed cells	c) Recombinant cells	d) None of these
199. The collection of bacteria	with gDNA is called:		
a) DNA clones		b) DNA library	
c) Genomic DNA library		d) cDNA library	
200. Which of the following st	atements are correct wi	th respect to a bioreactor?	
I. It can process small vol			
II. It provides optimum to			
일었다. 아프라스 전투하는 아이트 그리아 그리아 있다.	1. P. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.	pe reactor in which air is bul	bled
Choose the correct option			
a) I and II	b) I and II	c) II and III	d) I, II and III
201. PCR and Restriction Frag			
a) Genetic transformation		b) DNA sequencing	
c) Genetic fingerprinting		d) Study of enzymes	
202. Restriction enzymes may		128 9	
a) Making recombinant I		b) Gene mapping	
c) Diagnosis of genetic di		d) All the above	
203. Vent polymerase enzyme			
a) Thermococcus litorali	S	b) <i>Thermus aquaticus</i>	•
c) <i>E. coli</i>		d) Salmonella typhimur	
204. Genetically bacteria have			
a) Human insulin	b) Testosterone	c) Thyroxine	d) Melatonin
205. DNA fingerprinting meth		L) Faurusia atudisa	
a) DNA tests for identity	and relationships	b) Forensic studies	
c) Polymorphism	ale vanliaatina mini ahu	d) All of the above	
206. Plasmids are autonomou			a.i
a) Bacteriophage lambda	ļ	b) Leishmania donovar	
c) Escherichia coli	notoin in hactaria in a	d) Paramecium caudat	
207. Production of a human p	rotein in bacteria in gen	euc engineering is possible b	ecause:





a) Bacterial cell can carry out the RNA splicing reactions

b) The human chromosome can replicate in bacterial cell c) The mechanism of gene regulation is identical in humans and bacteria d) The genetic code is universal 208. Reverse transcriptase: a) Disintegrates host DNA b) Translates host DNA c) Transcribes viral RNA to DNA d) Polymerises host DNA 209. An example of gene therapy is: a) Production of injectable Hepatitis B vaccine b) Production of vaccines in food crops like potatoes which can be eaten c) Production of test tube babies by artificial insemination and implantation of fertilized eggs d) Introduction of gene for adenosine deaminase in persons suffering from Severe Combined Immuno-Deficiency (SCID) 210. Synthetic DNA or sDNA is: a) DNA synthesized in lab without any template b) DNA synthesized in the cell without any template c) DNA synthesized in the lab, without any nucleotide d) DNA synthesized in the cell without any nucleotide 211. Stirred-tank bioreactors have advantages over shake flasks because they a) Provide high temperature and pH b) Provide better aeration and mixing properties c) Do not allow the entry of CO₂ d) Are easy to operate 212. During 'gene cloning' which is called a gene taxi? b) Plasmid d) Protozoa a) Vaccine c) Bacteria 213. TATAATG sequence near the RNA start point of prokaryotic promoter is: a) Nicks b) DNA marker c) Pallindrome d) Pribnow box 214. I. Copy number is defined as the number of copies of plasmid present in a cell II. It varies from 15-100 copies per cell Choose regarding the above statements a) I is true, II is false b) II is true, I is false c) Both are true d) Both are false 215. Which one of the following hydrolyses internal phosphodiester bonds in a polynucleotide chain? b) Protease c) Exonuclease d) Endonuclease a) Lipase 216. What does Bt stand for the popular crop Bt cotton? a) Best b) Best type c) Biotechnology d) Bacillus thuringiensis 217. Which of the following statement is incorrect? a) Cosmid contains gene coding for viral protein b) Cosmid replicates like plasmids c) Cosmid has antibiotic resistant marker gene d) Cos site has 12 bases helping to join complete genome to make it circular 218. An attenuated virus: a) Is a virus that is non-pathogenic b) In an elongated viral particle c) Can transfer recombinant DNA to other viruses d) Will not produce an immune response 219. Which of the following has popularized the PCR (polymerase chain reaction)? a) Easy availability of DNA template b) Availability of synthetic primers c) Availability of cheap deoxyribonucleotides d) Availability of 'Thermostable' DNA polymerase 220. Choose the correct statement with reference to 'Dolly': a) She was created by taking nucleus from unfertilized eggs and cytoplasm from unfertilized eggs b) She was created by taking nucleus from under udder cells and cytoplasm from unfertilized eggs

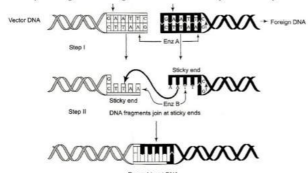


- c) She was created by taking cytoplasm from udder cell and nucleus from unfertilized eggs
- d) She was created by taking cytoplasm from udder cell and nucleus from fertilized eggs
- 221. The first recombinant DNA was constructed by
 - a) Stanley Cohen

b) Herbert Boyer

c) Both (a) and (b)

- d) Temin and Baltimore
- 222. Study the given diagram and identify the enzymes A and B involves in steps I and II



Step I

Step II

- a) *Eco* RI
- DNA ligase
- b) Alu I

DNA ligase

- c) Hind II
- DNA polymerase
- d) Restriction endonuclease DNA polymerase
- 223. Which one of the following is a correct statement
 - a) "Bt" in "Bt-cotton" indicates that it is a genetically modified organism produced through biotechnology
 - b) Somatic hybridization involves fusion of two complete plant cells carrying desired genes
 - c) The anticoagulant hirudin is being produced from transgenic Brassica napus seeds
 - d) "Flavr Savr" variety of tomato has enhanced the production of ethylene which improves its taste
- 224. The transgenic animals are those which have:
 - a) Foreign RNA in all its cell

- b) Foreign DNA in all its cells
- c) Foreign DNA in some of its cells
- d) Both 'A' and 'C'
- 225. Which of the following is not correctly matched for the organism and its cell wall degrading enzyme?
 - a) Plant cells-Cellulase
- b) Algae-Methylase
- c) Fungi-Chitinase
- d) Bacteria-Lysozyme
- 226. Petroleum-lysing bacteria are being engineering for the removal of oil spills. What is the most realistic danger of these bacteria to the environment?
 - a) Mutations leading to the production of a strain pathogenic to humans
 - b) Extinction of natural microbes due to the competitive advantage of the "petro-bacterium"
 - c) Destruction of natural oil deposits
 - d) Poisoning of the food chain
- 227. c-DNA probes are copied from the messenger RNA molecules with the help of:
 - a) Restriction enzymes

b) Reverse transcriptase

c) DNA polymerase

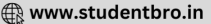
- d) Adenosine deaminase
- 228. Mishandling of genetic engineering may cause:
 - a) Genetic erosion
- b) Green revolution
- c) Silver revolution
- d) White revolution

- 229. Gene for cloning may be chemically synthesized:
 - a) When the exact sequence of nucleotides is known
 - b) Through the use of restriction enzymes and gel electrophoresis to separate restriction fragments
 - c) By the Sanger method
 - d) By making complementary DNA from genes without introns
- 230. Source of taq polymerase used in PCR is a
 - a) Thermophilic fungus

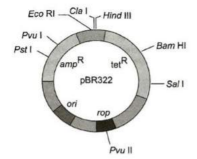
b) Mesophilic fungus

c) Thermophilic bacterium

- d) Halophilic bacterium
- 231. Genetic engineering has been successfully used for producing:
 - a) Transgenic models for studying new treatment for certain cardiac diseases



	b) Transgenic Cow-Rosie which produces high fat mi	lk for making ghee	
	c) Animals like bulls for farm work as they have supe	er power	
	d) Transgenic mice for testing safety of polio vaccine	before sue in humans	
232.	Which of the following is used as a best genetic vector	r in plants?	
	a) Bacillus thuringiensis	b) Agrobacterium fumefac	ciens
	c) Pseudomonas putida	d) None of the above	
233.	Plants in comparison to animals are more rapidly ma	nipulated by genetic engin	eering. Select out the most
	probable reason for this		
	a) Totipotency shown by plant cells		
	b) Single somatic cell can regenerate a whole plant b	ody	
	c) Genetic engineering is supplemented with plant ti	ssue culture techniques	
	d) All of the above		
234.	Which of the following pairs is correctly matched?		
	a) Central dogma-Codon	b) Okazaki fragments-Spli	cing
	c) RNA polymerase-RNA primer	d) Restriction enzymes-Go	enetic engineering
235.	Recombinant DNA technology is related with:		
	a) Stanley Cohen and Harbert Boyer	b) Bateson and Punnet	
	c) Huxley and Harvey	d) Schleiden and Schwann	1
236.	Western blotting technique was developed by:		
	a) Alwin b) Edwin	c) Towbin	d) Thomas
237.	In recombinant DNA technique, the term vector refer	rs to a	
	a) Donor DNA, it is identified and picked up through	electrophoresis	
	b) Plasmid, transfers DNA into living cell		
	c) Collection of entire genome in form of plasmid		
	d) Enzyme, cuts the DNA at specific sites		
238.	Complete transduction is:		
	a) Transfer of whole genome with the help of virus		
	b) Picking up of one or more genes by a phage and tr	ansfer it to second host	
	c) Integration of gene brought by viral particle into g	enome of new host	
	d) Both B and C		
239.	The function of polymerase chain reaction (PCR) is:		
	a) Translation b) Transduction	c) DNA amplification	d) None of these
240.	The steps involved in the Southern blot test are as fo	llows	
	I. X-ray film		
	II. Electrophoresis		
	III. Digestion with restriction enzyme		
	IV. Ethidium bromide		
	V. Radioactive probe		
	Choose the option having correct sequential order of	these events	
	a) III, II, IV, V and I b) III, IV, II, V and I	c) III, II, V, IV and I	d) II, IV, III, V and I
241.	The given figure is the diagrammatic representation	of the <i>E. coli</i> vector pBR32	2. Which one of the given
	options correctly identifies its certain component(s)	?	



a) Ori-original restriction enzymes

b) Rop-reduced osmotic pressure

c) Hind III, Eco RI-selectabel markers

d) amp^R, tet^R-antibiotic resistances genes

242. The restriction enzyme(s) used in recombinant DNA technology that make staggered cuts in DNA leaving sticky ends is/are

a) Eco RI

b) Hind II

c) Bam HI

d) All of the above

243. RNA processing is:

a) An event that occurs after RNA transcribed

b) The rejection of old, worn-out RNA

c) An event that occurs before RNA is transcribed

d) Both (A) and (C)

244. Find out the wrong statements

a) Mobile genetic elements, transposons were visualized by Barbara McClintock

b) Udder cell and somatic cell is used to produce the cloned sheep by nuclear transplantation method

c) In pedigree analysis, a person immediately affected by and action is called propositus

d) DNA ligases are used to cleave a DNA molecule

245. Widely used tool in genetic engineering of crop plants is:

a) Protoplast fusion

b) Transposon

c) Microinjection

d) Agrobacterium mediation

246. DNA fingerprinting method is very useful for:

a) DNA tests for identity and relationships

c) Polymorphism

b) Forensic studiesd) All of the above

247. Who among the following discovered the enzyme restriction endonuclease?

a) Hamilton Othanel Smith

b) Sir Godfrey Hounsfield

c) F. Jacob

d) Andre Lwoff

248. The mobile genetic element is

a) Transposons

b) Mutation

c) Endonuclease

d) Variation

249. The enzyme used for cutting DNA segment in genetic engineering is:

a) ATP-ase

b) Ligase

c) DNA polymerase

d) Restriction endonuclease

250. When the number of genes increases in response to some signal, the effect is called:

a) Gene dosage

b) Gene pool

c) Gene amplification

d) Gene frequency

251. Identify the palindromic sequence in the following

a) $\frac{GAATTC}{CTTUUG}$

b) $\frac{\text{GGATCC}}{\text{CCTAGG}}$

c) cc. 1 c

d) $\frac{CGATAC}{GCTAAG}$

252. Colony hybridization procedure for identification of plasmid clones is called:

a) Southern blotting

b) Grunstein-Hogness assay

c) DNA probes d) Molecular assay

253. The different basic steps of genetic engineering are given below randomly

I. Identification of DNA with desirable genes

II. Gene transfer

III. Maintenance of DNA in host and gene cloning

IV. Introduction of DNA into host to from recombinant DNA

Which of the following represents the correct sequence of steps?

a) I, II, III and IV

b) I, IV, III and II

c) III, IV, II and I

d) I, III, IV and II



- 254. Which of the following steps are involved in the process of recombinant biotechnology? Arrange in correct order
 - I. Extraction of the desired gene product
 - II. Amplification of the gene of interest
 - III. Isolation of a desired DNA fragment
 - IV. Ligation of the DNA fragment into a vector
 - V. Insertion of recombinant DNA into the host

Correct order is

- a) I, II, III, IV and V
- b) III, II, IV, V and I
- c) II, IV, V, III and I
- d) I, IV, V, III and II
- 255. In bacteria, genes for antibiotic resistance are usually located in:
 - a) Chromosomal DNA
- b) Cytoplasm
- c) Mitochondria
- d) Plasmids

- 256. Natural genetic engineer is:
 - a) Bacillus subtillis

b) Pseudomonas spp

c) Escherichia coli

- d) Agrobacterium tume faciens
- 257. A number of bacteria with recombinant DNA of same type form:
 - a) Clone library
- b) Gene library
- c) Gene pool
- d) Gene frequency

- 258. I. ...A... is the ability of a cell to take up foreign DNA
 - II. The cell is treated with specific concentration of a divalent cation such as ...B... to increase pore size in cell wall
 - III. InC... method recombinant DNA is directly injected into the nucleus of an animal cell

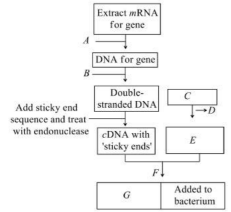
The most appropriate option regarding A, B and C is

- a) A-Competency, B-Calcium, C-gene gun method
- b) A-Transformation, B-Sodium, C-microinjection method
- c) A-Competency, B-Calcium, C-microinjection method
- d) A-Transformation, B-Sodium, C-gene gun method
- 259. T₁ plasmid is used for making transgenic plants. It is obtained from:
 - a) Azotobacter

b) Agrobacterium

c) Rhizobium in leguminous root

- d) Yeast
- 260. Identify and match the labelled items *A*, *B*, *C*, *D*, *E*, *F* and *G* in the diagram below from the list I-VII given with components



- I. DNA polymerase
- II. plasmid
- III. plasmid with 'sticky ends'
- IV. DNA ligase
- V. restriction endonuclease
- VI. recombinant DNA
- VII. reverse transcriptase
- The correct components are
- ABCDEFG

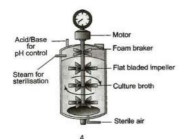


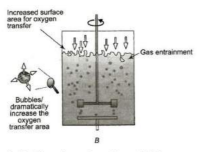


a) VII I II V III IV VI	b) VII VI V IV III II	
c) VII V III I II IV VI	d) I II IV VI III V V	
261. A technology which has found immense use in solving		
a) DNA fingerprinting	b) Polymerase chain read	
c) Recombinant DNA technology	d) Monoclonal antibody p	oroduction
262. The most important feature in a plasmid to be used		
a) Origin of replication	b) Presence of a selectab	le marker
c) Presence of sites for restriction endonuclease	d) Its size	N 2
263. DNA gyrase, the enzyme that participates in the pro-		type of
a) DNA ligase	b) DNA polymerase	
c) DNA topoisomerase	d) Reverse transcriptase	
264. Abnormal gene is replaced by normal gene through:		
a) Gene therapy b) Medicines	c) Cloning	d) Radiation
265. The key tools required for the recombinant DNA tec	hnology are	
I. restriction enzymes II. Polymerase enzymes		
III. host organism ligases IV. Vectors		
V. host organisms		
Select the correct option		
	c) I, II, III and V	
266. A tumour inducing plasmid widely used in the produ		
a) Escherichia coli	b) Bacillus thuringiensi	
c) Staphylococcus aureus	d) Agrobacterium tume	
267. Which one of the following palindromic base sequen	ices in DNA can be easily ci	ut at about the middle by
some particular restriction enzyme?		
a) 5'GATATG3'		
3'5'		
b) 5'GAATTC3'		
3'5'		
c) 5'CACGTA3'		
3'5'		
d) 5'CGTTCG3'		
3'5'		
268. Which of the following infection (s) can be diagnose	ranki san	chain reaction?
a) HIV-1 and HIV-2 viruses	b) Hepatitis-B virus	
c) Mycobacterium tuberculosis	d) All of the above	
269. Agarose is extracted from		
a) Sea weeds b) Blue-green algae	c) <i>Ephedra</i>	d) <i>Sargassam</i>
270. Which one is a true statement regarding DNA polym		
 a) It is used to ligate introduced DNA in recipient ce 	하게 되었다. 이 아이에서 하는 그래를 하네가 하네가 하셨다가 하게 하네가 있는데,	
c) It is isolated from a virus	d) It remains active at hig	5 - 5
271. DNA fragments generated by the restriction endonu		ion can be separated by:
a) Polymerase chain reaction	b) Electrophoresis	
c) Restriction mapping	d) Centrifugation	
272. The two main techniques that gave birth to modern	biotechnology are	
I. chemical engineering		
II. genetic engineering		
III. human genome engineering		
IV. molecular biology		
Choose the correct option		
a) I and II b) I and III	c) II and IV	d) II and III

273. Stirred-tank bioreactors have been designed for										
a) Purification of the product										
	b) Addition of preservatives to the product									
	c) Availability of oxygen throughout the process									
d) Ensuring anaerobic conditions in the culture vess										
274. First biochemical to be produced commercially by m		ic engineering is:								
a) Interferon b) Penicillin	c) Human insulin	d) Fertility factors								
275. Which is incorrect statement?										
 a) Taq DNA polymerase is important for PCR 										
b) Taq DNA polymerase is not thermostable										
c) In PCR two nucleotide primers are used										
d) Taq DNA polymerase, isolated from bacterium Th	AN OWNER OF THE PARTY CONTRACTOR OF THE PARTY CONTRACT									
276. A genetically engineered micro-organism used succe	5									
a) Trichoderma b) Xanthomonas	c) Bacillus	d) Pseudomonas								
277. There is a restriction endonuclease called <i>Eco</i> RI. Wh										
a) Coli b) Coelom	c) Coenzyme	d) Colon								
278. Which of the following would have the highest oxygo		stics?								
a) A sparged stirred tank bioreactor being stirred at										
b) A non-sparged stirred tank bioreactor being stirr	ed at 200 RPM									
c) A shake flask being mixed at 200 RPM										
d) All of the above would have equivalent oxygen tra										
279. Enzymes breaking nucleic acids into nucleotides are		20100 01								
a) Hydrolases b) Amylases	c) Nucleic acidases	d) Nucleases								
280. Palaeontologists unearthed a human skull during ex		and the second control of the second control								
attached to it. Only little DNA could be extracted from		ent man need to be analysed,								
the best way of getting sufficient amount of DNA fro	m this extract is									
a) By hybridizing the DNA with a DNA probe										
b) By subjecting the DNA to polymerase chain reacti	on									
c) By subjecting the DNA to gel electrophoresisd) By treating the DNA with restriction endonucleas										
281. Transgenic organisms are produced by:	es									
a) Deleting sex chromosomes	b) Inducing gene mutation	ne								
c) Introducing foreign genes	d) Arresting spindle fibre									
282. Manipulation of gene and genetic material by man is										
the formation of recombinant DNA molecules. This b										
a) Recombinant DNA technology	b) Genetic engineering	as								
c) DNA manipulation biotechnology	d) All of the above									
283. Ligases catalyse the formation of bonds between	a) An of the above									
a) $C = C$ b) $P = O$	c) C – C	d) H – H								
284. The characteristics of a molecular probe are	c) c c	dj II								
I. very long molecule										
II. double-stranded										
III. DNA or RNA										
IV. complementary to a part of desired gene										
The correct pair is										
a) I and II b) II and III	c) III and IV	d) IV and I								
285. VNTR analysis involves	.,	,								
a) Analyzing specific loci for two base repeating uni	ts usually less then 100 bp	in size								
b) Analyzing specific loci for 2-4 bp repeating units										
c) PCR amplification of specific genes										
a sa de										

d) Cutting DNA with restriction enzyme and analyzing	ng the banding pattern of fr	agments
286. Manipulation of DNA in genetic engineering became		
a) Restriction endonuclease	b) DNA ligase	,
c) Transcriptase	d) Primase	
287. Study the given figure carefully and select the correct		;
Wells	t statements regarding this	
DNA bands		
* A B		
I. It represents typical agarose gel electrophoresis w	hich showing differential n	nigration of DNA fragments
II. Lane 1 contains undigested DNA fragments		
III. Lanes 2 to 4 contains digested DNA fragment		
IV. Smallest DNA bands are present at (A) position a		present at (B) position
a) I, II and III b) I, II and IV	c) II and III	d) III and IV
288. Matching sequence of DNA between two evidences, of		Secretary Secretary
a) DNA fingerprinting b) DNA amplification	c) Gene mapping	d) DNA resolution
289. Alec Jeffreys developed the DNA fingerprinting techn	E	
a) Ribozyme b) Sex chromosomes	c) SNP	d) VNTR
290. In addition to <i>taq</i> polymerase enzyme which other t	hermostable DNA polymer	ases have been isolated to
be used in Polymerase Chain Reaction (PCR)?		
a) <i>Vent</i> polymerase b) <i>Pfu</i> polymerase	c) Both (a) and (b)	d) None of these
291. PCR proceeds in three distinct steps governed by ten		er of
a) Denaturation, synthesis (polymerization), anneali		
b) Annealing, synthesis (polymerization), denaturati		
c) Synthesis (polymerization), annealing, denaturati		
d) Denaturation, annealing, synthesis (polymerization	on)	
292. One of the following is transgenic of organisms:	13 77 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
a) Holly sheep and Flavr savr tomato	b) Holly sheep and Cottor	
c) Dolly sheep and Cotton Ct	d) Flavr savr tomato and	
293. Name of the drug used in cancer treatment produced		
a) HGH b) TSH 294. What is the function of Restriction endonuclease?	c) Insulin	d) Interferon
a) Restricts the synthesis of DNA inside the nucleusb) Synthesizes DNA		
c) Cuts the DNA molecule randomly		
d) Cuts the DNA molecule at specific sites		
295. I. Bacteriophages areA nfectectingB		
IIC are hybrid vectors derived from plasmids wh	nich contain or site of λ nha	nge
A, B and C in above statements refers to	nen contain of site of n pile	ige
A B C		
a) Protozoa Bacteria Cosmid	b) Plasmid Virus Co	smid
c) Bacteria Virus Cosmid	d) Virus Bacteria Cos	
296. In gel electrophoresis, the separated bands of DNA a		
is called		an and Ber breeze, thus seeb
a) Elution b) Origin replication	c) Competency	d) Transformation
297. Nif genes is a group of proteins:	way wassanan Parazanan A	voy voice are conserved to conserve the first
a) 15 genes b) 15 nucleotides	c) 15 proteins	d) 10 genes
298. Identify the following diagrams <i>A</i> and <i>B</i> and select the		(B) (100 B)
energen var mannet 🗲 programme på fatt sitt 🔀 uddat 🔀 supplement 1,500 fatt fra - kjeld 1,500 fra 1,500 fill 1,500 fil	er o region de propositione de Transporter	





- a) A-Simple stirred-tank bioreactor, B-Sparged stirred-tank bioreactor
- b) A-Sparged stirred-tank bioreactor, B-Complex stirred-tank bioreactor
- c) A-Sparged stirred-tank bioreactor, B-Simple stirred-tank bioreactor
- d) A-Simple stirred-tank bioreactor, B-Complex stirred-tank bioreactor
- 299. Genetic engineering is helpful in:
 - a) Gene regulation
- b) Gene translation
- c) Gene therapy
- d) Alcohol production
- 300. Significance of heat shock method in bacterial transformation is facilitate
 - a) Binding of DNA to the cell wall

- b) Uptake of DNA through membrane transport proteins
- c) Uptake of DNA through transient pores in the bacterial cell wall
- d) Expression of antibiotic resistance gene
- 301. A technique used to make numerous copies of a specific segment of DNA quickly and accurately:
 - a) Ligase chain reaction

b) Transcription

c) Polymerase chain reaction

- d) Translation
- 302. Two microbes found to be very useful in genetic engineering are:
 - a) Diplococcus sp. and Pseudomonas sp.
 - b) Crown gall bacterium and Caenorhabditis elegans
 - c) Escherichia coli and Agrobacterium tumefaciens
 - d) Vibrio cholerae and a tailed bacteriophage
- 303. Minisatellite or Variable Number Tendem Repeat (VNTR) are used in
 - a) Gene therapy
- b) Gene mapping
- c) DNA fingerprinting
- d) Restriction enzymes
- 304. Having become an expert on gel electrophoresis, you are asked to examine a gel for a colleague. Where would you find the smallest segment of DNA?
 - a) Near the positive electrode, farthest away from the wells
 - b) Near the negative electrode, close to the wells
 - c) Near the top, near the negative pole
 - d) Near the middle they tend to slow-down after the first few minutes
- 305. Improvement of genotype of an organism by addition of some foreign genes is:
- a) Genetic diversity
- b) Gene handling
- c) Tissue culture
- d) Genetic engineering

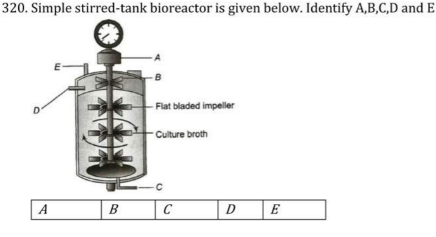
- 306. The structure involved in genetic engineering is
 - a) Codor
- b) Anticodon
- c) Vector
- d) Plasmid
- 307. In agarose gel electrophoresis, DNA molecules are separated on the basis of their
 - a) Charge only
- b) Size only
- c) Charge to size ratio
- d) All of these
- 308. In gel electrophoresis, the sample DNA is cut into fragments by







a) Restriction endonucleases b) Exonuclease c) Endonuclease d) Anhydro L-galactose 309. Molecular scissors, which cut DNA at specific site: a) Ligase b) Cellulase c) Pectinase d) Polymerase 310. PCR stands for: a) Polymerase Cyclic Reaction b) Polymerase Chain Reaction c) Polyethyl Cytosine Reaction d) Polymerization Chain Reaction 311. In case of polymerase chain reaction, temperature, required for the steps A. Denaturation B. Annealing C. Extension a) A-94°C, B-40°C, C-72°C b) A-40°C, B-72°C, C-94°C c) A-72°C, B-94°C, C-40°C d) A-94°C, B-72°C, C-40°C 312. DNA can be introduced into any cell by: a) Injection b) Being complexed with calcium salts c) Being placed along with the cell into a gene gun d) Gel electrophoresis 313. An improved variety of transgenic basmati rice: a) Gives high yield and is rich in Vitamin A b) Is completely resistant to all insect pests and diseases of paddy c) Gives high yield but has no characteristic aroma d) Does not require chemical fertilizers and growth hormones 314. Which of the following organelles is associated with genetic engineering? a) Plastids b) Plasmids c) Chloroplast d) Mitochondria 315. Human genome contains about: b) 10,000 genes a) 10,000 nucleotides c) 6 billion nucleotides d) 6 billion genes 316. An artificial process of infecting cells with naked viral DNA is: a) Translation b) Transduction c) Transfection d) Transgenic 317. Match the correct one: a) RNA Polymerase-RNA primer b) Respiration-Lysosome c) Restriction enzyme-genetic engineering d) Central dogma-DNA structure 318. For transformation, microparticles coated with DNA are to be bombarded with gene gun are made up of: a) Platinum or Zinc b) Silicon or Platinum c) Gold or Tungsten d) Silver or Platinum 319. You are attempting to introduce a gene that imparts larval moth resistance to bean plants. Which of the



following vectors are you most likely to use?

b) Bacterial plasmid

c) Ti plasmid

d) Yeast plasmid

a) Phage DNA

a)	Motor	Foam braker	Sterile air	Steam for sterilization	Acid/Bas e of pH control	b)	Foam braker	Sterile air	Steam for sterili	Acid/ Base of pH
				tale .	52 AS	3			zeation	control
c)	Acid/ Base of	Motor	Foam braker	Sterile air	Steam for sterilize	d)	Sterile air	Steam for	Foam braker	Motor

ation

321. Protein engineering is used to study the proteins to compare the catalytic properties of:

a) Normal and mutated form of enzyme

b) Normal form of enzyme

sterilize

ation

c) Mutated form of enzyme

d) Normal and mutated form of proteins

322. Genes that are involved in turning on or off the transcription of a set of structural genes are called:

a) Polymorphic genes

pH

control

b) Operator genes

c) Redundant genes

d) Regulatory genes

Acid/Bas

e of pH

control

323. The experimental manipulation of DNA of different species, producing recombination DNA is known as

a) Gel electrophoresis

b) Transformation

c) Genetic engineering

d) Replication technology

324. Plasmid is used as carrier because:

a) It has both ends with replicating points

b) It has no free ends

c) It is circular DNA with a capacity of binding with equkaryotic DNA

d) All of the above

325. Which of the following statement is correct in the context of observing DNA separated by agarose gel electrophoresis?

a) DNA can be seen in visible light

b) DNA can be seen without staining in visible light

c) Ethidium bromide stained DNA can be seen in visible light

d) Ethidium bromide stained DNA can be seen under exposure to UV light

326. Nitrogen fixing genes are called:

a) 'Nif' genes

b) Plasmid genes

c) Leg genes

d) Cos genes

327. The genetically-modified (GM) brinjal in India has been developed for:

a) Enhancing shelf life

b) Enhancing mineral content

c) Drought-resistance

d) Insect-resistance

328. Variable number of tendem repeats (VTNRs) in the DNA molecule are highly useful in:

a) Monoclonal antibody production

b) DNA fingerprinting

c) Recombinant DNA technology

d) Stem cell culture

329. Protoplasts of two different species are fused in:

a) Clona propagation

b) Organography

c) Micropropagation

d) Somatic hybridization

330. Identify the correct match for the given diagram

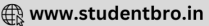


Apparatus function

a) Gene gun - Vectorless direct gene transfer

b) Electrophoresis - Differential migration of DNA fragments





c) Bioreactor - Raw materials are biologically converted into specific products

d) Respirometer - Finding out rate of respiration

331. DNA fingerprinting technique was first developed by:

a) Jeffreys, Wilson and Thein

b) Schleiden and Schwann

c) Edward and Steptoe

d) Boysen and Jensen

332. Using recombinant technology, genes from a donor cell can be transplanted into a bacterium for DNA replication and protein synthesis. The kinds of cells that can be used as a donor in this technology are

a) Bacteria

b) Either yeast or bacteria

c) Eukaryotic cells

d) Any kind of cell

333. Transformation is defined as the procedure by which a piece of ...A... is introduced into a ...B... host. Here A and B refers to

A B

a) RNA Virus

b) DNA Bacteria

c) RNA Bacteria

d) DNA Virus



BIOTECHNOLOGY PRINCIPLES AND PROCESSES

						: ANS	W)	ER K	EY:						
1)	a	2)	b	3)	c	4)	c	169)	c	170)	b	171)	d	172)	(
5)	a	6)	b	7)	a	8)	a	173)	b	174)	a	175)	a	176)	ä
9)	C	10)	C	11)	C	12)	d	177)	d	178)	a	179)	C	180)	(
13)	C	14)	C	15)	c	16)	a	181)	d	182)	a	183)	b	184)	é
17)	d	18)	d	19)	a	20)	b	185)	a	186)	C	187)	a	188)	(
21)	b	22)	a	23)	d	24)	255	189)	d	190)	C	191)	c	192)	é
25)	b	26)	a	27)	b	28)	d	193)	a	194)	C	195)	b	196)	ä
29)	a	30)	a	31)	a	32)	d	197)	d	198)	b	199)	a	200)	(
33)	d	34)	a	35)	C	36)	a	201)	d	202)	d	203)	a	204)	ć
37)	d	38)	d	39)	a	40)	d	205)	d	206)	C	207)	d	208)	(
41)	d	42)	a	43)	c	44)	b	209)	d	210)	a	211)	b	212)	ł
45)	b	46)	a	47)	b	48)	c	213)	d	214)	C	215)	d	216)	(
49)	a	50)	d	51)	a	52)	a	217)	a	218)	a	219)	d	220)	ł
53)	b	54)	c	55)	d	56)	c	221)	C	222)	a	223)	C	224)	ł
57)	C	58)	d	59)	b	60)	C	225)	b	226)	C	227)	d	228)	ć
61)	b	62)	C	63)	d	64)	С	229)	a	230)	C	231)	d	232)	ł
65)	C	66)	a	67)	c	68)	c	233)	d	234)	b	235)	a	236)	(
69)	b	70)	a	71)	a	72)	d	237)	b	238)	C	239)	c	240)	ć
73)	a	74)	a	75)	b	76)	a	241)	d	242)	d	243)	a	244)	(
77)	a	78)	d	79)	d	80)	d	245)	d	246)	d	247)	a	248)	í
81)	b	82)	c	83)	a	84)	d	249)	b	250)	C	251)	b	252)	ł
85)	a	86)	b	87)	a	88)	b	253)	b	254)	b	255)	d	256)	•
89)	a	90)	d	91)	d	92)	a	257)	b	258)	C	259)	b	260)	ä
93)	b	94)	d	95)	a	96)	a	261)	a	262)	a	263)	C	264)	ä
97)	b	98)	C	99)	d	100)	d	265)	d	266)	d	267)	b	268)	(
101)	d	102)	c	103)	d	104)	c	269)	a	270)	d	271)	b	272)	ä
105)	b	106)	a	107)	b	108)	b	273)	c	274)	C	275)	b	276)	(
109)	b	110)	b	111)	b	112)	b	277)	a	278)	a	279)	d	280)	ł
113)	d	114)	d	115)	c	116)	c	281)	c	282)	d	283)	b	284)	(
117)	a	118)	C	119)	c	120)	a	285)	d	286)	a	287)	a	288)	ä
121)	a	122)	c	123)	b	124)	d	289)	d	290)	C	291)	d	292)	(
125)	a	126)	a	127)	a	128)	b	293)	d	294)	d	295)	C	296)	ä
129)	b	130)	a	131)	d	132)	d	297)	a	298)	a	299)	c	300)	(
133)	b	134)	d	135)	a	136)	d	301)	c	302)	C	303)	c	304)	é
137)	d	138)	c	139)	d	140)	a	305)	d	306)	d	307)	b	308)	ä
141)	a	142)	c	143)	b	144)	b	309)	c	310)	b	311)	a	312)	ł
145)	d	146)	c	147)	c	148)	a	313)	a	314)	b	315)	c	316)	(
149)	c	150)	b	151)	c	152)	b	317)	c	318)	c	319)	c	320)	ä
153)	b	154)	c	155)	a	156)	a	321)	a	322)	b	323)	c	324)	(
157)	b	158)	a	159)	d	160)	d	325)	d	326)	a	327)	d	328)	ŀ
161)	d	162)	c	163)	c	164)		329)	d	330)	c	331)	a	332)	(
165)	c	166)	a	167)	a	168)		333)	b	*** **********************************				neddette man i	

BIOTECHNOLOGY PRINCIPLES AND PROCESSES

: HINTS AND SOLUTIONS :

2 **(b)**

Retroviruses in animals including humans are able to change normal cells into cancerous cell

4 (c)

pBR322 vector was the first artificial cloning vector constructed in 1977 by Boliver and Rodriquez. It is widely used in gene cloning experiments in pBR322

p - Denotes that it is plasmid

BR – stands for Boliver and Rodriquez who constructed this plasmid

322 is a number given to distinguish this plasmid from others developed in the same laboratory

5 (a)

Genetic engineering is defined as the modification of genetic information of living organism by direct manipulation of their DNA. Thus, a gene of known function (economic importance) can be transferred from its normal location into a cell *via* a suitable mobile genetic element called vector such as plasmid, phage, etc.

7 (a)

Recombinant DNA having integrated fragment of antibiotic resistant gene

8 (a)

True. In plants, the tumour inducing plasmid (T_i) of $\mbox{\it Agrobacterium tume faciens}$ is used as a cloning vector

9 (c)

Gene encoding resistance to antibiotics like ampicillin, chloramphenicol, tetracycline or Kanamycin, are useful selectable markers for *E.coli*. The normal *E.coli* cells do not carry resistance against any of these antibiotics

12 (d)

Proteins are removed by treatment with protease

13 (c)

Plasmids, cosmids or bacteriophages can be used as vector in genetic engineering. Plasmids are most widely used circular, extrachromosomal DNA segments seen in the bacterial cells. They carry a foreign gene or desired gene to the host.

The size of plasmids ranges from 1×10^6 to 200×10^6 daltons

14 (c)

Both are true, *Ori* also controls the copy numbers of the linked DNA

If a foreign DNA ligates at the *Bam* HI site tetracycline resistance gene in the vector pBR322, the recombinant plasmid loses the tetracycline

18 (d

After the formation of the product in the bioreactors, it undergoes through some processes before a finished product to be ready for marketing. *The processes include* (i) separation and (ii) purification of product which are collectively called the downstream processing The product is subjected to quality control testing and kept in suitable preservatives. If drugs are to be manufactured such formulation has to undergo through clinical trials. A proper quality control testing for each product is also needed. The downstream processing and quality control test are different from product to product

19 (a)

Endonucleases are enzymes that produce internal cuts called cleavage DNA molecule. A class of endonucleases cleavage DNA only within or near those sites which have specific base sequences, such endonucleases are known as restriction endonucleases and sites recognized by them are called recognition sites. Restriction endonucleases have major role in genetic engineering

20 **(b)**

Plasmid is an extrachromosomal genetic of DNA that is capable of replicating independently of host chromosome. It forms the basis of many cloning vectors used in genetic engineering

21 (b)

PCR was discovered by Kary Mullis. In Polymerase Chain Reaction (PCR), a segment of DNA is amplified. *Taq* DNA polymerase enzyme is used PCR, this enzyme is temperature resistant



22 (a)

A-Taq polymerase, B-Denaturation (air), C-Prime

23 (d)

Bioreactors (fermenters) are considered as vessel in which raw material are biologically converted into specific products by microbes, plant and animal cells and/or their enzymes

24 (a)

By using PCR phenylketonuria, muscular cystrophy, sickle-cell anaemia, hepatitis, chlamydia and tuberculosis can be diagnosed

26 (a)

Primers are small chemically synthesized oligonucleotides of about 10-18 nucleotides long that are complementary to the sequences present at the 3' ends of the target DNA segment

27 **(b)**

Shotgun cloning involves cutting the DNA of the entire genome into pieces with restriction enzyme, inserting these pieces or fragments into bacteria or yeast with plasmids or viruses and allowing the organism to reproduce making copies or clones of the DNA fragments

28 (d)

The Polymerase Chain Reaction or PCR, as it is commonly called, was originally invented by Kary Mulllis in 1985. Kary Mulllis shared the Nobel Prize with Michael Smith in Chemistry in 1993. PCR is best defined as the DNA replication *in vitro*. A single PCR amplification cycle involves three basis steps; denaturation, annealing and extension (polymerization)

30 (a)

True, *Ori* is a DNA sequence that is responsible for initiating replication. Any piece of DNA, which linked to this sequence can replicated with in the host cells

31 (a)

True. Plasmids are autonomously replicating circular extra-chromosomal DNA

33 (d)

PCR is carried out in the following three steps Denaturation, Annealing and Extension

37 **(d)**

Plasmid which is extra chromosomal DNA molecule and help in gene cloning

38 (d)

A restriction fragment containing a specific gene of interest can be identified by gel electrophoresis followed by transferring of DNA to a membrane as a solid support matrix using a procedure called a Southern blot

39 (a)

Protection of host DNA from the action of restriction endonuclease by adding methyl group to one or two bases usually with in the sequence recognized by restriction enzyme

40 (d)

Single stranded DNA molecules that can bind to and be used to detect other DNA molecule are called probes

42 (a)

Principle of PCR The purpose of a PCR (Polymerase Chain Reaction) is to make a huge number of copies of a gene. This is necessary to have enough starting template for sequencing There are three major steps in a PCR, which are repeated for 30 or 40 cycles. This is done on an automated cyclers, which can heat and cool the tubes with the reaction mixture in a very short time

- (i) **Denaturation at 95**°C During the denaturation, the double-strand melts open to single-stranded DNA, all enzymatic reactions stop (for example : the extension from a previous cycle)
- (ii) Annealing at 54°C The primers are jiggling around, caused by the Brownian motion. Ionic bonds are constantly formed and broken between the single-stranded primer and the single-stranded template. The more stable bounds last a little bit longer (primers that fit exactly) and on that little piece of double-stranded DNA (template and primer), the polymerase can attach and starts copying the template. Once there are a few bases built in, the ionic bond is so strong between the template and the primer, that it does not break anymore
- (iii) Extension at 72°C This is the ideal working temperature for the polymerase. The primers, where there are a few bases built in, a already have a stronger ionic attraction to the template than the forces breaking these attractions. Primers, that are on positions with no exact match, get loose again (because of the higher temperature) and don't give an extension of the fragament

The bases (complementary to the template) are coupled to the primer on the 3' side (the polymerase adds *d*NTPs from 5' to 3', reading the

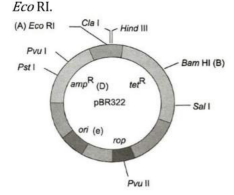


template from 3' to 5' side, bases are added complementary to the template)

43 **(c)**

The controlled use of biological agents, such as microorganism, plants or animal cell, for beneficial use is called biotechnology. This technology involves cutting and pasting of desired DNA fragments into the specified hosts for the benefits of human beings

51 (a)



52 (a)

Microinjection DNA is directly injected into plant protoplasts or cells (specifically into the nucleus or cytoplasm) using fine tipped (0.5-1.0 micrometer diameter) glass needle or micropipette. This method of gene transfer is used to introduce DNA into large cells such as oocytes, eggs, and the cells of early embryo

Electroporation It involves a pulse of high voltage applied to protoplasts/cells/tissues to make transient (temporory) pores in the plasma membranes which facilitates the uptake of foreign DNA

The cells are place in a solution containing DNA and subjected to electrical shock to cause holes in the membranes. The foreign DNA fragments enter through the holes into the cytoplasm and then to nucleus

Chemical Mediated Gene Transfer Chemicals like Polyethylene Glycol (PEG) and sulphate induce DNA uptake into plant protoplasts. Calcium phosphate is also used to transfer DNA into cultured cells

55 (d)

Polyethylene glycol method is used for gene transfer without a vector. It is a chemical method for direct gene transfer to protoplast

56 **(c)**

Restriction endonucleases and ligase are commonly used enzymes in genetic engineeering

57 (c

DNA fingerprinting is a modern technique that compares sets of DNA by locating identical sequences of nucleotides. It is oftening used to solve many mysteries involving murders, robberies and rapes

58 (d)

Genetic engineering is a branch of biotechnology, which deals with the manipulation of genetic material by man. The technique of genetic engineering includes

- (i) formation of 'recombinant DNA'
- (ii) use of gene cloning
- (iii) gene transfer
- pBR 322 was the first artificial cloning vector constructed in 1977 by Boliver and Rodriguer. It is widely used in gene cloning experiments
- 2. Restriction enzymes belongs to a class of enzymes called nucleases

60 **(c)**

A - Key Mullis

B - 1985

C-1993

61 (b)

Cutting of piece of DNA from a plasmid was done with the help of restriction enzyme, popularly known as molecular scissors

62 (c)

Different kinds of specific enzymes are used in genetic engineering, *e.g.*, cleaving enzymes → These enzymes are used to break DNA molecules *They are of three types*

- (i) Exonucleases
- (ii) Endonucleases
- (iii) Restriction endonucleases

63 (d)

Components of a bioreactors

An agitator system

An oxygen delivery system

Foam control system

Temperature control system

pH control system

sampling ports to withdraw culture periodically

- 65 **(c)**Both are true
- 66 (a)







A-plasmid, B-Boliver, C-Rodriquez. pBR322 vector was the first artificial cloning vector constructed in 1977 by Boliver and Rodriquez. It is widely used in gene cloning experiments in pBR322

p - Denotes that it is plasmid
 BR - stands for Boliver and Rodriquez who constructed this plasmid
 322 is a number given to distinguish this plasmid from others developed in the same laboratory

67 (c)

DNA fingerprinting is a technique to identify a person on the basis of person's DNA specificity. The technique is based upon the fact that the DNA constitution of an individual carries some specific sequence of nucleotides, which do not carry any information for protein synthesis

From the given options, leucocytes are to be used for identifying the criminal because they are nucleotide, whereas erythrocytes are enucleated

70 **(a)**

The basic requirements of a PCR reaction are the following

DNA Template Any source that contains one or more target DNA molecules to be amplified can be taken as template

Two Nucleotide Primers Primers, which are oligonucleotides, that hybridise to the target DNA region, one to each strand of the double helix **Enzyme** *Taq* polymerase and *vent* polymerase

72 **(d)**

Circular plasmid DNA which is used as a vector, can be cleaved at one site with the help of enzyme to give a linear DNA molecule. A foreign DNA segment can now be inserted, by joining the ends of broken circular DNA to the two ends of foreign DNA, thus regenerating a bigger circular DNA molecule that can now be separated by gel electrophoresis on the basis of its size Bacteriophages provide another source of cloning vectors. Since, usually, a phage has a linear DNA molecule, a single break will generate two fragments, which are later joined together with foreign DNA to generate a chimeric phage particle

73 (a)

Genetic engineering is defined as the modification of genetic information of living organisms by direct manipulation of their DNA

Thus, a gene of known function (or economic importance) can be transferred from its normal

location into a cell *via* a suitable mobile genetic element called vector such as plasmid phage, etc.

74 (a

Thermostable enzymes 'Taq and Vent' isolated from thermophilic bacteria are DNA polymerase Taq polymerase, isolated from a Thermophilic bacterium, Thermus aquaticus and vent polymerase, isolated from a thermophilic bacterium Thermococcus litoralis

75 (b

Due to chlorophenicol resistance gene, one is able to select a transformed cell in the presence of chloramphenicol. The chloramphenicol resistance gene in this case is called selectable marker

76 (a)

The restriction endonuclease *Eco* RI is obtained from *Esherichia coli* RY 13. The recognition sequence for this is GAATTC, CTTAAG

77 (a)

Autonomously replicating circular extrachromosomal DNA.

Manipulation of gene and genetic material by man is a fast emerging branch of science, which started with the formation of recombinant DNA molecule. This branch of science is named as recombinant DNA technology, genetic engineering and DNA manipulation technology, genetic engineering and DNA manipulation technology. This technology involves cutting and pasting of desired DNA fragments into the specified hosts for the benefits of human beings

78 (d)

The polymerase chain reaction is a technique that is used for *in vitro* replication of specific DNA sequence using thermostable DNA polymerase. The polymerase chain reaction or PCR, was originally invented by Kary Mullis in 1985. Kary Mullis shared the Nobel Prize with Michael Smith in chemistry in 1993

86 **(b)**

The Polymerase Chain Reaction (PCR) is a technique by which small samples of DNA can be quickly amplified. The repeated amplification is achieved by the use of thermostable DNA polymerase (*i.e., taq* polymerase isolated from a bacterium, *Thermus aquaticus*) which remain active during the high temperature induced denaturation of double-stranded DNA

88 (b)





Exonucleases remove nucleotides from the terminal ends (either 5' or 3') of DNA in one strand of duplex

90 (d)

PCR is a technique of synthesizing multiple copies of the desired gene or (DNA) *in vitro. The basic requirement of PCR* are DNA template, two nucleotide primers and enzyme (DNA polymerase)

91 (d)

Agrobacterium tumefaciens (soil inhabiting plant bacterium) is a pathogen of several dicot plants. It delivers a piece of DNA known as 'T-DNA' in the Ti-plasmid which transforms normal plant cells into tumour cells to produce chemicals against pathogens

92 (a)

Restriction endonuclease recognize a specific DNA base sequence (recognition sequence, recognisation site, restriction sequence or restriction site having palindromic sequence) and cleaves both the strands of DNA at or near that site. The enzyme cuts the DNA, generating restriction fragments with overhanging ends or blunt ends

95 (a)

Agrobacterium tumefaciens (updated scientific name Rhozobium radiobacter) is the casual agent of crown gall disease (the formation on tumour) in over 140 species of dicot. It is a rod-shaped, Gram negative, soil bacterium (Smith, et. al 1907). Symptoms are caused by the insertion of a small segment of DNA, known as T-DNA (transfer DNA) into the plant cell, which is incorporated at a semi-random location into the plant genome

96 (a)

True, the polymerase chain reaction is a reaction in which amplification of specific DNA sequences is carried out in *vitro*

99 (d)

Restriction enzyme are known as molecular knives or molecular scissors and are used to cut DNA at specific sites of DNA. These were first discovered by Smith, Nathan and Arber

101 (d)

Small volume cultures are usually employed in laboratories for research and production of less quantities of products. *e.g.,* in shake flasks. However, large scale production of the products is carried out in 'bioreactor'

Bioreactors are large vessels (having a volume of 100 to 1000 L) which are used for biological conversion of raw materials into specific products. The most commonly used bioreactors are of stirring type

102 (c)

The term 'Biotechnology' was given in 1917 by a Hungarian Engineer, Karl Erkey, to describe a process or large scale production of pigs

107 (b)

Agrobacterium tumefaciens delivers a piece of DNA known as 'T-DNA' in the Ti-plasmid which transforms normal plant cells into tumour cells to produce chemical against pathogens

110 **(b)**

Kary Mullis

Gene encoding resistance to antibiotics like ampicillin, chloramphenicol, tetracycline or Kanamycin, are useful selectable markers for *E.coli*. The normal *E.coli* cells do not carry resistance against any of these antibiotics

114 (d)

Ti-plasmid is found in *Agrobacterium* tumefaciens, which produces crown gall (tomour) in a large number of dicot species. *A. tumefaciens* is a Gram negative soil bacterium that infects a wide range of plants and causes crown galls

115 (c)

The science of recombinant technology took birth when Cohen and Boyer (1972) were able to introduce a piece of antibiotic resistance gene containing foreign DNA into plasmid of *Salmonella typhimurium*. This modified plasmid was them inserted into *E. coli* to get clones of recombinant DNA. Thus, Cohen and Boyer discovered recombinant technology

116 (c)

In recombinant DNA technology, a desired segment of DNA or a gene is made to combine with the DNA of an organism where it will multiply and produce it copies. Plasmids and viruses are the most commonly used cloning vectors in recombinant DNA technology

119 (c)

Selectable marker helps to select the host cells which contain the vector and eliminate the non-transformants. Genes encoding resistance to antibiotics like ampicillin, chloramphenicol, tetracycline or kanamycin are useful selectable





markers of E.coli. The normal E.coli cells do not carry resistance against any of these antibiotics

122 (c)

Herbert Boyer discovered that restriction enzymes have the capability of cutting DNA strands in a particular fashion, which left what has became known as sticky ends on the strands

123 (b)

A Southern blot.

A restriction fragment containing a specific gene of interest can be identified by gel electrophoresis 139 (d) followed by transferring of DNA to a membrane as a solid support matrix using a procedure called a Southern blot

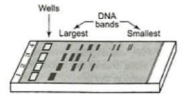
124 (d)

In biolistic or gene gun method, cells are a high velocity micro-particles of gold or tungsten coated with DNA in plants. Important crop plants like maize, rice and wheat have now been transformed by this method

125 (a)

Electrophoresis.

A molecule of DNA can be cut into fragments by the enzyme restriction endonucleases. These fragments of DNA can be separated by a technique of gel electrophoresis. In this process the smallest | 142 (c) segment of DNA travel towards anode (+ ve electrode), farthest away from the wells



130 (a)

RNA is removed by treatment with ribonuclease

All statements are correct

Restriction Enzymes	Source	Recognition Sequence and Site of Cleavage	Product
Eco RI	Escherichia coli RY 13	5'-G-A-A-T-T-C-3' 3'-C-T-T-A-A-G-5'	G A-A-T-T-C C-T-T-A-A G Sticky ends

133 (b)

During annealing two oligonucleotide primers hybridise to each of single stranded template DNA in presence of excess of synthetic oligonucleotides

136 (d)

In gel electrophoresis a molecule of DNA can be cut into fragments by the enzyme restriction

endonuclease. DNA fragments move towards the anode according to their molecular size through the agarose gel

The separated DNA fragments can be observed only after staining them with a solution of ethidium bromide. The bright orange coloured bands of DNA can be seen only under UV light. These bands of DNA fragments are cut out from the gel and extracted by using convenient technique. This step is called elution

Microorganisms can be grown in the bioreactors by support growth system and suspended growth

141 (a)

Escherichia coli and Agrobacterium tumefaciens are the microbes found to be very useful in genetic engineering. E.coli is a motile, Gram negative, rod-shaped bacterium which is a normal inhabitant of human colon. It is most extensively used in bacterial genetic and molecular biology Agrobacterium tumefaciens is a soil bacterium. It has Ti-plasmid (tumour inducing plasmid) and it can be used for the transfer of a desired gene in dicot plants

pUC 18 is a plasmid cloning vector commonly used with E. coli. The vector length is 2686 bp and is isolated from *E. coli* strain DH5α by standard procedures

143 (b)

A - Vector; B-DNA

144 (b)

The probes used for DNA fingerprinting are usually prepared from minisatellite or microsatellite DNA

145 (d)

In recent times, PCR is being used in the detection of HIV (virus of AIDS) mutation are related to genetic disease. By using PCR phenylketonuria, muscular dystrophy, sickle-cell anaemia, hepatitis, chlamydia and tuberculosis can be diagnosed. PCR is also used in DNA fingerprinting

147 (c)

Ti-plasmid is a plasmid present in Agrobacterium tumefaciens. It is used in genetic engineering in plants, e.g., as a vector in gene transfer to dicot plants

148 (a)





The role of DNA ligase in the construction of a recombinant DNA molecule is formation of phosphodiester bond between two DNA fragments. DNA ligase help in sealing gaps in DNA

Therefore, they act as a molecular glue. In 1969 Har Govind Khorana discovered DNA ligase in T₄bacteriophage

153 **(b)**

In gene gun or biolistic method tungsten or gold particles, coated with foreign DNA are bombarded into target cells at a very high velocity Although this method is suitable for plants yet this technique is also used to insert genes into animal that promote tissue repair into cells (particularly cancer of mouth) near wounds

154 (c)

The final step in PCR is extension (polymerization), where in Taq DNA polymerase synthesizes the DNA region between the primers using deoxynucleotide triphosphates and Mg²⁺. It | 163 (c) means the primers are extended towards each other so that the DNA segment lying between the two primer is copied. The optimum temperature for this polymerization step is 72°C Taq polymerase is thermostable enzyme, isolated from Thermophilic bacterium, Thermus aquaticus

155 (a)

EFB - European Federation of Biotechnology A definition of biotechnology which covers both traditional views and modern molecular biotechnology has been given by European Federation of Biotechnology. According to EFB "Biotechnology is the integrated use of biochemistry, microbiology and engineering sciences in order to achieve technological application of the capabilities of microorganisms, cultured tissues/cells and part there of"

156 (a)

A technique developed by EM Southern in 1975 for detection of a specific DNA sequences (gene or other) in a large, complex sample of DNA (e.g., cellular DNA). It is also used to determine the molecular weight of a restriction fragment and to measure relative amounts in different sample **Uses** Southern blots are used in gene discovery and mapping, evolution and development studies, diagnostics and forensics In regards to genetically modified organisms,

Southern blotting is used as a definitive test to

ensure that a particular section of DNA of known genetic sequence has been successfully incorporated into the genome of the host organism

157 (b)

Cry I endotoxins obtained from Bacillus thuringiensis are effective against bollworm larvae

158 (a)

In the naming of restriction enzymes the first letter is derived from genus name and next two letters from the species name of the prokaryotic cell from where the enzymes are extracted

159 (d)

A molecule of DNA can be cut into fragments by the enzyme restriction endonucleases. These fragments of DNA can be separated by a technique of gel electrophoresis. It is a technique used for the separation of substances of different ionic properties

During extension, the enzymes Taq polymerase synthesizes the DNA segment between the primers. The two primers extend towards each other in order to copy the DNA segment typing between the two primers This step requires presence of deoxynucleoside triphosphate (dNTPs) and Mg²⁺ and occurs at 72°C

164 (c)

Both are true in the process for the isolation of DNA, after several treatments the purified DNA is precipitated by adding chilled ethanol. The bacterial/plant, animal cell is broken down by enzymes to release DNA, along with RNA, proteins, polysaccharide and lipids

165 (c)

Bioreactors are vessels of large volumes (100-1000 litres) in which raw materials are biologically converted into specific products. It provides all the optimal conditions for achieving the desired product by providing optimal growth conditions like temperature, pH, substrate, salts vitamins and oxygen. Stirred-tank bioreactors are commonly used bioreactors. There are cylindrical with curved base to facilitate proper mixing of the contents. The stirrer mixes the contents and makes oxygen available throughout the bioreactor

166 (a)

Thermus aquaticus.





DNA polymerase which is stable at high temperature (>90°C) is required to carry out the synthesis of new DNA. The DNA polymerase like *Taq* polymerase is generally used in PCR reactions which is isolated from a bacterium *Thermus aquaticus*

169 (c)

The first restriction endonuclease type II was isolated by Smith, Wilcox and Kelley from *Haemophilus influenza* bacterium. It was formed to cut DNA molecules at a particular point of recognizing a specific sequence of six base pairs, known as the recognition sequence

170 (b)

In gel electrophoresis, the separated DNA fragments are visualized after staining the DNA with ethidium bromide followed by exposure to UV radiation

173 (b)

In gel electrophoresis a molecule of DNA can be cut into fragments by the enzyme restriction endonuclease. DNA fragments move towards the anode according to their molecular size through the agarose gel

The separated DNA fragments can be observed only after staining them with a solution of ethidium bromide. The bright orange coloured bands of DNA can be seen only under UV light. These bands of DNA fragments are cut out from the gel and extracted by using convenient technique. This step is called elution

175 (a)

DNA polymerase which is stable at high temperature (>90°C) is required to carry out the synthesis of new DNA. The DNA polymerase like *Taq* polymerase is generally used in PCR reactions which is isolated from a bacterium *Thermus aquaticus*

176 (a)

Most sensitive technique to detect malignant cell in non-hodgkins lymphoma is polymerase chain reaction. In recent times, PCR is being used in the detection of HIV (Virus of AIDS)

179 (c)

The Pribnow box (also known as the Pribnow – Schaller box) is the sequence TATAAT of six nucleotides that is an essential part of a promoter site on DNA for transcription to occur in bacteria

187 (a)

Gene gun method was first developed by Prof. Stanford and coworkers at Cornell University, USA in 1987. This method is used to introduce foreign DNA into host cell

188 (c)

During extension, the enzyme DNA polymerase synthesizes the DNA segment between the primers. DNA polymerase is a heat stable enzyme

191 (c)

After the formation of the product in the bioreactors, it undergoes through some processes before a finished product to be ready for marketing. The processes include (i) separation and (ii) purification of products, which are collectively called the downstream processing

192 (a)

The stirred-tank bioreactor is well suited for large-scale production of protein of enzyme by using microbial plant/animal/human cells

193 (a)

A-DNA is vector/plasmid DNA and B-is foreign DNA.

C-The restriction enzyme that recognizes this palindrome-*Eco* RI

D-The enzyme that can link these two DNA fragment-DNA ligase

194 (c)

Restriction endonuclease was isolated for the first time by W Arber in 1962 in bacteria. They are called molecular scissors or biological scissors. In 1978 Arber, Smith and Nathan were awarded the Nobal Prize for the discovery of restriction endonuclease

195 (b)

In genetic engineering *r*DNA technology is applied to several biotechnological processes for obtaining particular biochemical improvement of genetic make up of an organism and fighting genetic defects

197 (d)

Primer and DNA polymerase.

PCR is a technique of synthesizing multiple copies of the desired gene or (DNA) *in vitro. The basic requirement of PCR* are DNA template, two nucleotide primers and enzyme (DNA polymerase)

198 (b)

An antibiotics resistance gene in a vector usually helps in the selection of transformed cell

200 (c)



Bioreactors are considered as vessels in which raw materials are biologically converted into specific products by microbes, plant and animal cells and or their enzymes. Small volume cultures can not give large quantities of the products. Large scale production (100-1000 L) of the products is carried out in bioreactors. A bioreactor provides the optimal conditions for obtaining the desired product by providing optimum growth conditions such as temperature, pH, substrate, vitamins, oxygen and salts. In the sparged stirred tank bioreactor, sterile air bubbles are sparged. The surface area for oxygen transfer is increased

203 (a)

Vent polymerase enzyme used in PCR is isolated from *Thermococcus litoralis*

211 (b)

A stirred-tank bioreactor is more advantageous, than shake flasks. It has an agitator system to mix the contents properly, an oxygen delivery system to make availability of oxygen, a foam control system, a temperature control system, a pH control system and a sampling port to withdraw the small volumes of the culture periodically

212 (b)

During gene cloning plasmid is called gene taxi. Molecular biologists add desired gene desired gene to plasmids, then insert the new plasmid with the added gene into a living bacterium

214 (c)

Both are true. Copy number is defined as the number of copies of vectors present in a cell. It varies from 1-100 copies per cell

219 (d)

Availability of thermostable DNA polymerase. DNA polymerase which is stable at high temperature (>90°C) is required to carry out the synthesis of new DNA. The DNA polymerase like *Taq* polymerase is generally used in PCR reactions which is isolated from a bacterium *Thermus aquaticus*

221 (c)

Stanley Cohen and Herbert Boyer generated first recombinant DNA molecule by combining a gene from a bacterium with plasmid of *Escherichia coli*

230 (c)

Thermophilic bacterium.

Thermostable enzymes '*Taq* and *Vent*'isolated from thermophilic bacteria are DNA polymerase

Taq polymerase, isolated from a Thermophilic bacterium, Thermus aquaticus and vent polymerase, isolated from a thermophilic bacterium Thermococcus litoralis

232 (b)

Agrobacterium tumefaciens is used as a best genetic vector in plants

233 (d)

Plants in comparison to animals are more rapidly manipulated by genetic engineering reasons are
(i) Totipotency (having the ability to differentiate

- into all cell types) shown by plant cells
 (ii) Single somatic cell can regenerate a whole
- plant body
 (iii) Genetic engineering is supplemented with
 plant tissue culture techniques

237 (b)

Vector is a plasmid or virus DNA used to introduce genes into a host cell, where the genes may be amplified (gene cloning) or otherwise manipulated

240 (a)

Digestion with restriction enzyme

33

Electrophoresis

1

Ethidium bromide

1

Radioactive probe

1

X-ray film

241 (d)

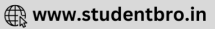
 amp^R (amplification resistance gene) and tet^R (tetracycline resistance gene) are antibiotic resistance genes

244 (d)

Restriction endonucleases cleave DNA molecules only at specific nucleotide sequence called restriction sites. DNA ligase enzymes is used to joins bits of DNA

248 (a)

Mobile genetic element is broadly any genetic element capable of moving itself, with or without duplication, from one site in a genome to another. Mobile genetic elements include plasmids, viruses, transposable genetic elements (transposons), short interspread elements, pathogenicity islands and so on. The term 'transposon' was introduced RW Hedges and AE Jacob in 1974, 'controlling elements' or jumping



genes, discovered by **Barbara McClintock** (1950) in maize

251 **(b)**

Special sequence in the DNA recognized by restriction endonuclease is called palindromic nucleotide sequence.

Restriction endonuclease recognizes palindromic sequences in DNA and cuts them

The palindromes in DNA are base pair sequences that are the same when read forward (left to right) or backward (right to left) from a central axis of symmetry

For example

253 (b)

Identification of DNA with desirable gene

Introduction of DNA into host to form recombinant DNA



Maintenance of DNA in host and gene cloning

↓ •------

Gene transfer

254 **(b)**

Recombinant DNA technology involved the following steps

- (i) Isolation of DNA
- (ii) Fragmentation of DNA by restriction endonucleases
- (iii) Isolation of a desired DNA fragment
- (iv) Amplification of the gene of interest
- (v) Ligation of the DNA fragment into a vector
- (vi) Insertion of recombinant DNA into the host
- (vii) Culturing the host cells on a suitable medium at a large scale
- (viii) Extraction of the desired gene product
- (ix) Downstream processing of the products as finished product, ready for marketing

258 (c)

- A Competency
- B Calcium
- C microinjection method

262 (a)

The most important feature in a plasmid to be used as a vector is origin of replication (*Ori*). Origin of replication is a specific sequence of DNA bases which is responsible for initiating

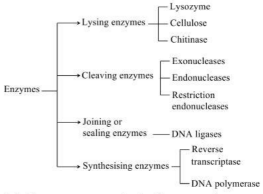
replication. A prokaryotic DNA has a single origin of replication while eukaryotic DNA may have more than one origin of replication

263 (c)

DNA gyrase, the enzyme that participates in the process of DNA replication is a type of DNA topoisomerase

265 (d)

Three types of 'biological tool' are used in the formation of recombinant DNA



- (ii) Cloning vectors (vehicle vectors)
- (iii)Complementary host (for transformation with recombinant DNA)

268 (d)

In recent times PCR is being used in the detection of HIV (Virus of AIDS). By using PCR phenylketonuria, muscular dystrophy, sickle-cell anaemia, hepatitis, chlamydia and tuberculosis also can be diagnosed

269 (a)

Agarose is extracted from sea weeds. It is a polysaccharide. In gel electrophoresis, DNA fragments separate according to size through the pores of agarose gel

270 (d)

DNA polymerase remains active at high temperature. Usually *Taq* DNA polymerase, isolated from a thermophilic bacterium *Thermus aquaticus*, is used in most of the cases

272 (a)

The science of biotechnology is based mainly on two core technologies

- (i) Genetic engineering, which is the manipulation of genes by man. It includes techniques to alter the nature of genetic material (DNA and RNA), to introduce these into host organisms and thus, change the phenotype of the host organism
- (ii) **Biochemical engineering**, *i.e.*, processes that help the growth of desired microbe/eukaryotic cell in large quantities in a sterile medium for the



manufacture and multiplication of biotechnological product

273 (c)

Each bioreactor has a cylindrical stirred-tank to facilitate the mixing of contents. The stirrer provides facility of mixing the contents as well as availability of oxygen throughout the process

275 (b)

Taq DNA polymerase is a thermostable enzyme, isolated from a *Thermophilic bacterium, Thermus aquaticus*

278 (a)

A sparged stirred-tank bioreactor being stirred at 200 RPM

280 (b)

The Polymerase Chain Reaction (PCR) is a technique by which small samples DNA can be quickly amplified. Starting with only one gene sized pieces of DNA, this technique is used to make literally billions of copies in only a few hours

283 (b)

Ligase catalyse the formation of bonds between P = 0

284 (c)

A probe is radioactively labelled (P³²) nucleic acid (20-40 nucleotide long) with a short sequence complementary to at least one part of the desired DNA gene

285 (d)

VNTRs were an important sources of RFLP genetic markers used in linkage analysis of genomes. VNTRs have become essential to forensic crime investigations, *via* DNA fingerprinting

286 (a)

Isolation of restriction endonucleases by **Nathans** and **Smith** (1970) made it possible to cut DNA at specific sites. Restriction enzyme can cut both strains of DNA when foreign nucleotides are introduced in the cell. They cleave DNA to generate a nick with a 5' phosphoryl and 3' hydroxyl terminus

287 (a)

Largest DNA bands will be at (A) and smallest DNA bands will be at (B) because in this DNA is move according to their size in agarose small DNA fragment will have small resistant so this fragment move to long distance as compared to large DNA fragment

289 (d)

The technique of fingerprinting was initially developed by Alec Jeffreys. He used a satellite DNA as probe that shows very high degree of polymorphism. It was called as Variable Number of Tendem Repeats (VNTRs)

290 (c)

Vent polymerase and pfu polymerase both

291 (d)

A single PCR amplification cycle involves three basic steps; denaturation, annealing and extension (polymerization) Denaturation – Melting of target DNA Annealing – Join

295 (c)

A-Bacteria, B-Virus, C-Cosmid

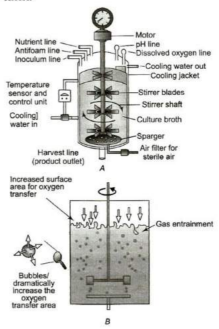
Extension - Polymerisation

296 (a)

The DNA fragments are seen as orange coloured bands. The separated bands of DNA are cut out and extracted from the gel piece. This step is called elution

298 (a)

Simple stirred-tank bioreactor, sparged stirred-tank.



A-Simple stirred-tank bioreactor for continuous culture.

B-Sparged stirred-tank bioreactor through, which sterile (free from any germs) air bubbles are sparged

Bioreactor (fermenters) Bioreactors are considered as vessels in which raw materials are biologically converted into specific products by







microbes, plant and animal cells and/or their enzymes. Small volume cultures cannot give large quantities of the products. Large scale production (100-1000 L) of the products in carried out in bioreactors. A bioreactor provides the optimal 307 (b) condition for obtaining the desired product by providing optimum growth conditions such as temperature, Ph, substrate, vitamins, oxygen and salts

Types of Bioreactors The most commonly used bioreactors are of **stirring type**. Stirring type bioreactors are (i) Simple stirredtank bioreactors and (ii) Sparged stirred-tank bioreactor as shown in figure. In the sparged stirred-tank bioreactor, sterile air bubbles are sparged. The surface area for oxygen transfer is increased

300 (c)

DNA being a hydrophilic molecule can not pass through cell membranes. Therefore, the bacteria should be made competent to accept the DNA

In this case the cell is treated with specific concentration of a divalent cation such as calcium to increase pore size in cell wall

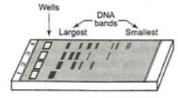
The cells are incubated with recombinant DNA on ice, followed by placing them briefly at 42°C and them putting it back on ice. This is called heat shock treatment. The bacteria now takes up the recombinant DNA

303 (c)

DNA fingerprinting technique is very useful in solving disputed parentage cases and forensic cases. DNA fingerprinting are obtained from RFLP and VNTR (satellite DNA) analysis of blood, hair or other material found the place of crime

304 (a)

A molecule of DNA can be cut into fragments by the enzyme restriction endonucleases. These fragments of DNA can be separated by a technique of gel electrophoresis. In this process the smallest segment of DNA travel towards anode (+ ve electrode), farthest away from the wells



306 (d)

The structure involved in genetic engineering is plasmid. Plasmids were discovered by William

Hays and Joshua Lederberg (1952). These are extrachromosomal, self-replicating usually circular, double-stranded DNA molecules found naturally in many bacteria and also in some yeast

After the cutting of DNA by restriction enzymes fragments of DNA are formed. Separation of DNA fragments according to their size or length is done by a technique called gel electrophoresis developed by A Tiselius in 1937

308 (a)

In gel electrophoresis, the sample DNA is cut into fragments by restriction endonucleases

311 (a)

A-Denaturation - 94°C B-Annealing - $40^{\circ} - 60^{\circ}$ C C-Extension - 72°C

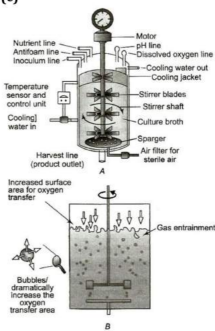
323 (c)

Genetic engineering

325 (d)

The separated DNA fragments can be seen only after staining the DNA with a compound known as ethidium bromide (E + Br) followed by exposure to UV radiation as bright orange coloured bands

330 (c)



A-Simple stirred-tank bioreactor for continuous

B-Sparged stirred-tank bioreactor through, which sterile (free from any germs) air bubbles are sparged

Bioreactor (fermenters) are Bioreactors considered as vessels in which raw materials are





biologically converted into specific products by microbes, plant and animal cells and/or their enzymes. Small volume cultures cannot give large quantities of the products. Large scale production (100-1000 L) of the products in carried out in bioreactors. A bioreactor provides the optimal 332 (d) condition for obtaining the desired product by providing optimum growth conditions such as temperature, Ph, substrate, vitamins, oxygen and 333 (b) salts

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A variety of cell types are used as a donor in recombinant DNA technology

A-DNA; B-Bacteria

