MARKING SCHEME (2023-24)

Class XII

Biotechnology (Subject Code-045)

Biotechnology (Subject Code-043)				
Q. No.	Answer	Marks		
	Section - A			
I	(b) They lyse specifically within the restriction site.	I		
2	(d) Prion	I		
3	(a) It measures both live and dead cells.	I		
4	(c) Alkaline Phosphatase	I		
5	(d) All of these	I		
6	(c) HeLa cell line	I		
7	(d) Whey	I		
8	(d) Diosgenin	I		
9	(b) To identify protein networks in nuclear pore complex.	I		
10	(d) SV40	I		
11	(c) Cystic Fibrosis	I		
12	(a) Dextran	I		
13	(a) Both Assertion and Reason are true and reason is the correct explanation of the assertion.	I		
14	(c) Assertion is true but the reason is false.	I		
15	(b) Both the Assertion and reason are true but reason is not the correct explanation of the assertion.	I		
16	(c) The assertion is true but the reason is false.	I		
	Section – B			
17	X is Subtilisin. The native enzyme subtilisin is easily inactivated by bleach (up to 90%). Solution to the problem is to use the detergent that contains Subtilisin that is modified by Site directed mutagenesis which is not affected by bleach. $(1/2x4=2)$	2		
18	Safety for human or animal consumption/ Effect on Biodiversity/Effect on beneficial insects or microbes Gene pollution/Development of superweeds/Change in fundamental vegetable nature of plants/ Antibiotic resistance in humans or animal pathogens/Changes in evolutionary pattern. (Any 4 for ½ mark each)	2		

19	Preparation is time consuming/Required one preparation to another. Trypsin is used to dissociate the adhered by the second secon		(Any two for ½ m		2	
	OR					
	Finite Cell Lines		Continuous Cell Lines			
	grow upto a limited number of generations		Grow continuously			
	Finite cell lines show contact inhibition, density limitation and anchorage dependence		No contact inhibition and an dependence. Density limitation reduced.	ichorage lost or		
	/ Finite cell lines show slow growth doubling time as 24-96 hours	rate or	continuous cell lines show rapid with doubling time as 12 to 24 hou	•		
		(A	Any two points of difference with 1 m	ark each)		
20	FISH	Karyo	typing		2	
	Interphase chromosomes can be used	Metaphase chromosomes are needed		(1 Mark)		
	Easy Technique as it gives colour to the chromosome	No su	ch specific colour	(1 Mark)		
21	(a) Protein samples A and B will get separated using this set up. (1 Mark)			(1 Mark)	2	
	(b) Using ampholytes with broader ran able to isolate all the four proteins.	nge cov	ering pH value range from 3 to 11 v	vill be (1Mark)		
		Sect	ion – C			
22	Replica plating.				3	
	Plasmid pBR322 carrying the insert in tet ^r gene in Multiple Cloning Sites (MCS) is used to transform the host cells which are first plated on solid media containing ampicillin. Overnight colonies from every single cell plated will develop which all have the plasmid. Replica plating is next performed to select colonies from this plate which are tetracycline sensitive due to insertional inactivation. The non recombinant colonies will grow on media with tetracycline and thus differentiate between recombinant and non recombinant cells.					
23	In Situ Activation means activation of zymogens at their site of activity in the presence of their biological target by alteration in its shape. (1 Mark)				3	
	Due to constellation of three amino acids because of unique folding of chymotrypsin, the asp 102 is able to hydrogen bond with the adjacent his 57 by borrowing a hydrogen ion. The his 57 in turn attracts a hydrogen ion from the adjacent ser 195 which allows its negatively charged oxygen anion to be able to make a nucleophilic attack on the peptide bond of the substrate. (2 Marks)					
24	(a) Lab media contain highly purifie economically used for large scale		•	can't be	3	
	(b) Provides uniform mixing of the medium and avoids development of anaerobic pockets thus ensuring optimum oxygen availability for growth.					
	(c) Foaming denatures the proteins	so it is u	indesirable. (1 x	3 marks)		

	OR	
	Somaclones through tissue culture, Mutant selection where mutants are produced using a mutagen like UV light, or Genetic Engineering can improve the production of the active compound.	
	(Any 2 for 1 Mark each)	
	The gene can be put under the control of a regulatory switch such that the production of recombinant protein does not occur until required. (1 Mark)	
25	The name of the technique is Protoplast Fusion and chemicals fusion like PEG can be used to fuse protoplasts from two different plants/ Electro-fusion. (1 Mark)	3
	Somatic hybrids and Cybrids can be produced using this method. (1 Mark)	
	Example: Intergeneric somatic hybrid between potato and tomato called Pomato/Topato or inter specifc somatic hybrid between two species of <u>Nicotiana</u> (any one, 1Mark)	
26	(a) Introduction of modified gene that encodes for overproduction herbicide target enzyme into crop plant making it insensitive to herbicide.	3
	(b) Introduction of gene that encodes for Bt toxin into the crop plant.	
	(c) Introduction of gene that encodes for viral coat protein into the crop plant. (1 x 3Marks)	
27	Leukemia, Heart disease/Heart attack, Paralysis/Spinal cord injury, Alzheimer's disease, Parkinson's disease, Huntington's disease, Burns	3
	(Any 6 for ½ mark each)	
28	(a) rHuEPO is used to treat anemia due to kidney failure/cancer treatment/treatment of AIDS/ blood loss during surgery. (Any one for 1 Mark)	3
	(b) tPA is used for dissolution of blood clots during a heart attack or stroke. (1 Mark)	
	(c)OKT3 binds to CD3 receptors of T lymphocytes causing immuno-suppression thus preventing rejection of kidney transplant. (1 Mark)	
	Section – D	
29	(i) 16 DNA molecules would be generated after 4 cycles. (1 Mark)	4
	(ii) Both the strands will act as the template in this case. (1 Mark)	
	(iii) 5' CTGAA 3' and 5' CAATT 3' (2 Marks)	
	OR	
	(iii)PCR can amplify the genome sequence from parents and offspring and DNA fingerprinting can match the pattern obtained. (2 Marks)	
30	 (i) Metabolite specific purification methods used are solvent extraction/ ion exchange chromatography/ salt precipitation. (Any two for ½ Mark each) 	4
	(ii) Flocculation/ Centrifugation/Ultrafiltration. (Any two for ½ Mark each)	
	(iii) For higher yields/higher stability of proteins/ cost reduction. (Any two for 1 Mark each) OR	
	(iii) Using specific Antibodies and probes which enable the detection of the organism	
	capable of producing specific products. (2 Marks)	

	Section-E		
31	(a) Restriction site of EcoRI is 5'-GAATTC-3'	(1 Mark)	5
	The ends generated will be called sticky. No, all the Restriction sequences may not be palindromic.	(½ Mark) (½ Mark)	
	(b)Microinjection can inject foreign DNA into plant and animal cells		
	Biolistics makes use of particle gun to bombard gold coated DNA Into c		
	(c)Small size of vector facilitates entry of recombinant molecules into the ho	. ,	
		o, 1 Mark each)	
	OR		
	(a)3' AGCTTCAGTC 3'	(1 Mark)	
	(b)Principle – When a ddNTP gets incorporated in the growing chain, the due to non availability of 3'hydroxyl group.	e reaction stops (1 Mark)	
	Steps- Each test tube out of four carries single stranded DNA templar DNA polymerase. Small amount of four ddNTPs are added separately i tubes. For example in test tube containing ddATP, all chains will termi at different positions of T present in the template. The prematu fragments are resolved and read with agarose gel electrophoresis. Marks)	nto the four test nate at ddA but	
32	Steps of Protein Fingerprinting	(5 Marks)	5
	Purify Haemoglobin Sickle cell RBC		
	B Trypsin treatment scHemoglobin Hemoglobin Hemoglobin is cleaved into small peptides by protease trypsin. Trypsin breaks peptide bonds adjacent to a lysine or an argining. scHemoglobin		
	Paper Electrophoresis		
	Paper chromatography		
	 Result : All peptides were similar from both samples except one (marked blue). 		
	Peptide sequencing		
	Protein fingerprinting		
	OR		

