

Program and Course Structure

School of Engineering Technology

M. Tech – Biotechnology

Program code: SET0203

Batch: 2018-20



1. Standard Structure of the Program at University Level

1.1 Vision, Mission and Core Values of the University

Vision of the University

To serve the society by being a global University of higher learning in pursuit of academic excellence, innovation and nurturing entrepreneurship.

Mission of the University

- 1. Transformative educational experience
- 2. Enrichment by educational initiatives that encourage global outlook
- 3. Develop research, support disruptive innovations and accelerate entrepreneurship
- 4. Seeking beyond boundaries

Core Values

- Integrity
- Leadership
- Diversity
- Community



Vision of the School

To become a globally acclaimed institution of higher learning in engineering and technology promoting excellence in research, innovation and entrepreneurship to provide sustainable solution to the needs of the society

Mission of the School

- 1. To impart quality education with strong industry & academic connectivity in the expanding fields of Engineering and Technology in a conducive and enriching learning environment.
- 2. To produce technocrats equipped with technical & soft skills and experiential learning required to stay current with the modern tools in emerging technologies to fulfill professional responsibilities and uphold ethical values.
- 3. To inculcate a culture of interdisciplinary research, innovation and entrepreneurship to provide sustainable solutions to meet the growing challenges and societal needs.
- 4. To foster collaborative learning and to play adaptive leadership role in professional career and pursuit of higher education through effective mentoring and counseling.

1.2.1Vision and Mission of the Department

Vision of the Department

To serve the society by being a global centre of higher learning in pursuit of academic excellence, innovation and nurturing entrepreneurship to cater to the needs of biotechnology in health, agriculture and environment sectors.

Mission of the Department

- M1: To conduct cutting edge multidisciplinary original research in plant, animal, medical, industrial and environmental biotechnology.
- M2: To train and transform students into thinking bioengineers, and scientists who are able to integrate theoretical knowledge with practical applications in diverse areas of Biotechnology
- M3: To adapt and update with rapidly changing technologies through self improvement with continuous learning and education, without compromising with moral and professional ethics.
- M4: To provide opportunities for collaborative learning beyond classrooms, in the broader community across the diverse spectrum of disciplines.



1.3 Program Educational Objectives (PEO)

1.3.1 Writing Program Educational Objectives (PEO)

The Educational Objectives of PG Program in Animal Biotechnology are:

- PEO1: Post Graduates will be able to integrate the biological sciences with engineering principles for the study of biological systems and medical health related problems.
- PEO2: Post Graduates will demonstrate the applications of bioengineering principles through development of industrial designs and processes that are of societal and industrial importance.
- PEO3: Post Graduates will update their knowledge and skill set with recent discoveries through self-improvement, research experience and continuous learning to create engineering solutions for society and environment.
- PEO4: Post Graduates will develop communication skills and demonstrate independent thinking, analytical and problem solving skills, self-management and function effectively in team oriented and open ended activities in an industrial or academic environment.
- PEO5: Post Graduates will develop leadership skills at levels appropriate to their experience and perform ethically and professionally in business, academia, industry and society.

Methods of Forming PEO's

- STEP 1: The needs of the Nation and society are identified through scientific publications, industry interaction and media.
- STEP 2. Taking the above into consideration, the PEOs are established by the Coordination Committee of the department.
- STEP 3. The PEOs are communicated to the alumni and their suggestions are obtained.
- STEP 4. The PEOs are communicated to all the faculty members of the department and their feedback is obtained.
- STEP 5. The PEOs are then put to the Board of Studies of the department for final approval.



1.3.3 Program Outcomes (PO's)

- PO1: **Engineering knowledge**: Apply the knowledge of engineering fundamentals, biological and physical sciences to the solution of complex engineering problems.
- PO2: **Problem analysis**: Identify and analyze complex engineering problems, formulate research solutions and reach substantiated conclusions using principles of basic and applied sciences and related technologies.
- PO3: **Design/development of solutions**: Design solutions for complex engineering problems and design system components or processes that meet the specified needs with appropriate consideration for the public health and safety, and the cultural, societal, and environmental considerations.
- PO4: **Conduct investigations of complex problems**: Use researchbased knowledge and research methods including design of experiments, analysis and interpretation of data, and synthesis of the information to provide valid conclusions.
- PO5: **Modern tool usage**: Create, select, and apply appropriate techniques, resources, and modern engineering and bioinformatics tools including prediction and modelling to study complex biological systems with an understanding of the limitations.
- PO6: **The engineer and society**: Apply reasoning informed by the contextual knowledge to assess societal, health, safety, legal and cultural issues and the consequent responsibilities relevant to the professional engineering practice.
- PO7: **Environment and sustainability**: Understand the impact of the professional engineering solutions in societal and environmental contexts, and demonstrate the knowledge of, and need for sustainable development.
- PO8: **Ethics**: Apply ethical principles and commit to professional ethics and responsibilities and norms of the engineering practice.
- PO9: **Individual and team work**: Function effectively as an individual, and as a member or leader in diverse teams, and in multidisciplinary settings.
- PO10: **Communication**: Communicate effectively on complex engineering activities with the engineering community and with society at large, such as, being able to comprehend and write effective reports and design documentation, make effective presentations, and give and receive clear instructions.
- PO11: **Project management and finance**: Demonstrate knowledge and understanding of the engineering and management principles and apply these to one's own work, as a member and leader in a team, to manage projects and in multidisciplinary environments.
- PO12: **Lifelong learning**: Recognize the need for, and have the preparation and ability to engage in independent and lifelong learning in the broadest context of technological change.



- PSO1: Acquire practical knowledge of animal system and related techniques to study life processes and apply the knowledge for research and industrial applications.
- PSO2: Ability to unravel metabolic and molecular pathways in animal cells and harness or manipulate them for human health and industrial products.
- PSO3: Develop understanding of recent research events through self learning and awareness in biotechnology and apply the acquired concepts for industrial purpose.
- PSO4: Conduct safe research and learn sustainable product development without compromising environmental safety and ethics.



1.3.5 The components of the curriculum

Course Component	Curriculum Content (% of total number of credits of the program)	Total number of contact hours	Total number of credits
Basic Sciences	3.75%	6	6
Engineering Sciences	9.06%	22	14.5
Humanities and Social sciences	3.12%	5	5
Technical and communications skills	10%	29	16
Sciences	13.4%	26	21.5
Program Core	27.5%	51	44
Program Electives	13.1%	21	21
Open Electives	6.8%	11	11
Project(s)	13.1%	36	21



School of Engineering and Technology M. Tech in Biotechnology

Batch: 2018-20 TERM: I

S.	Course	Course		eaching	Load	Credits		
No.	Code		L T P			Credits		
THEORY CLASSES								
1.	BTY601	Analytical Instruments for	3	1	0	4		
		Biotechnology						
2.		Elective1	3	0	0	3		
3.		Elective2	3	3 0 0		3		
4.		Elective3	3 1 0		4			
5.	BTY605	Molecular Cell Biology 3 0 0				3		
PRAC	CTICALS					•		
6.	BTP615	Enzyme & Genetic Engineering Lab	0	0	4	2		
7.	7. BTP605 Molecular Cell Biology Lab. 0 0 4					2		
	TOTAL CREDITS							



School of Engineering and Technology M Tech in Biotechnology

Batch: 2018-20 TERM: II

S.		Subjects	Tea	ching L	oad			
No.	Subject	L T P		P	Credits			
	Code							
THE	THEORY CLASSES							
1	BTY613 Biological database and their management 3 0 0		0	3				
2		Elective4	3	1	0	4		
3		Elective5	3	1	0	4		
4		Elective6 3 0 (0	3			
5		Elective7	Elective7 3 0 0		0	3		
6	MRM001	Research Methodology (MOOC)	Research Methodology (MOOC) 2 0 0			2		
PRA	CTICALS							
7	BTP606	Applied Bioinformatics Lab.	0	0	2	1		
8	BTP630	Cell and Tissue Engineering Lab.		0	2	1		
9	9 CCU101 Community Connect		0	0	4	2		
				T	OTAL	23		



School of Engineering and Technology M Tech in Biotechnology Batch: 2018-20

TERM: III

			oad	Credits		
	${f L}$	T	P	Credits		
PRACTICALS						
8 Seminar	0	0	4	2		
20 Dissertation I	0	0	20	10		
		Т	OTAL	12		
51		Seminar 0	518 Seminar 0 0 520 Dissertation I 0 0	518 Seminar 0 0 4		



School of Engineering and Technology M Tech in Biotechnology

Batch: 2018-20 TERM: IV

S.			Teaching Load			Credita
No.	Code		L	T	P	Credits
PRAC	CTICALS					
1	BTP621	Dissertation II	0	0	32	16
				T	OTAL	16



Syllabus

Prepared by Department of Biotechnology



Core

BTY601 Analytical Instruments for Biotechnology

Scl	hool: SET	Batch: 2018-2020
Pro	ogram: M	Current Academic Year: 2018-19
Te	ch	
Br	anch:	Semester: 01
Bio	otechnology	
1	Course Code	BTY601
2	Course Title	Analytical Instruments for Biotechnology
3	Credits	4
4	Contact	310
	Hours	
	(LTP)	
	Course Status	Compulsory
5	Course	To develop and understanding of the principle, instrumentation, operation
	Objective	and applications of different analytical, separation and diagnostic
		techniques used in the fields of Biochemistry, Molecular Biology and
		Biotechnology.
6	Course	CO1: Perform experiments based on electrophoretic techniques for
	Outcomes	separating proteins and nucleic acids.
		CO2: Purify compounds from a mixture using column, ionexchange,
		affinity chromatography, HPLC, affinity and gas chromatography.
		CO3: Apply the spectroscopy techniques (Absorption and fluorescence,
		atomic and circular dichroism) to characterize physiochemical
		properties of biological molecules. Determine structure and mass of
		organic compounds and proteins by nuclear magnetic resonance (NMR),
		mass spectrometry and Xray crystallography.
		CO4: Review imaging techniques for disease diagnosis.
		CO5: Illustrate organelle and protein localization by microscopy. Isolate
		cells by using fluorescence activated cell sorting (FACS) and magnetic
		activated cell sorting (MACS). Purify proteins by ultrafiltration and
		dialysis for enzymatic reactions and protein blotting.
		CO6: Relate the basic instrumentation techniques with practical
<u> </u>	~	applications for Biotechnology.
7	Course	This course will cover the major topics on electrophoretic techniques for
	Description	separating proteins and nucleic acids, purify compounds from a mixture
		using column, ionexchange, affinity chromatography, HPLC, affinity and
		gas chromatography, spectroscopy techniques (Absorption and
		fluorescence, atomic and circular dichroism) to characterize physiochemical



Outline syllabu	compounds spectrometry diagnosis, r sorting (FAC by ultrafiltra relate the base Biotechnolo	and proteins and Xray and Xray nicroscopy, ICS) and magration and dialastic instrume	nolecules, determine structure and mass of organic s by nuclear magnetic resonance (NMR), mass crystallography, imaging techniques for disease solate cells by using fluorescence activated cell netic activated cell sorting (MACS), purify proteins lysis for enzymatic reactions and protein blotting, entation techniques with practical applications for			
•		rocic				
			sis (Southern, Northern and Western blotting)			
			ctrophoresis: Principle and applications			
			1 11			
		•	merpic and applications			
			chromatography			
		Raman spectroscopy and NMR: Instrumentation and working				
			-			
			and fluorescence, Atomic spectroscopy), Xray			
		• •	1 1			
Unit 4		crystallography: crystal preparation, working and uses Medical Imaging and Spectrometry				
			ring of mass spectrometry			
	-		ument setup and working of FACS			
C			<u> </u>			
Mode of						
examination		Theory, and fit factions with				
Weightage	CA	MTE	ETE			
Distribution	30%	20%	50%			
Text book/s*	Wilson K. and Walker J., "Principles and Techniques of Biochemistry and Molecular Biology", Cambridge University Press, 2010.					
Other			D.P. and Benore M., "Fundamental Laboratory			
References		•	nemistry and Biotechnology", Wiley, 2009.			
		-	cal Biochemistry: Principles and Applications",			
	Unit 1 A B C Unit 2 A B C Unit 3 A B C Unit 4 A B C Unit 5 A B C Unit 5 A B C Unit 5 A B C T C Unit 5 A B C O C Unit 5 A B C O C O C C C C C C C C C C C C C C C	compounds spectrometry diagnosis, resorting (FAG by ultrafiltrarelate the basiotechnolo) Outline syllabus Unit 1 Electrophon A Principle of B Capillary and C 2Dgel electry Unit 2 Chromatog A Column and B Affinity and C HPLC: Instruction Unit 3 Spectroscopy A Raman spectory B Spectrophoto C Spectroscopy crystallograpy Unit 4 Medical Im A Magnetic References A Optical, AFI B Ultracentrific C Ultrafiltration Mode of Theory/Jury examination Weightage CA Distribution 30% Text book/s* Wilson K. and Molecular B Other References Approace 2. Sheehan	compounds and proteins spectrometry and Xray diagnosis, microscopy, I sorting (FACS) and magre by ultrafiltration and diagnosis, microscopy, I sorting (FACS) and magre by ultrafiltration and diagnosis, microscopy. Outline syllabus Unit 1			



BTY 605: Molecular Cell Biology

	hool: School of	Batch: 2018-20				
	gineering &					
	chnology					
	ogram: M.Tech	Current Academic Year: 2018-19				
	anch:	Semester: 1				
Bie	otechnology					
1	Course Code	BTY 605				
2	Course Title	Molecular Cell Biology				
3	Credits	3				
4	Contact Hours (LTP)	300				
	Course Status	Compulsory				
5	Course Objective	On successful completion of this module students will be able to:				
		 Determine the role of different types of channels associated with trafficking of the molecules. Predict the translocation of biomolecules between different cell 				
		organelles 3. Visualize cells and cellular organelles using microscopy.				
		4. Analyze metabolic activities of a cell and the production of metabolic energy in form of ATP				
		5. Characterize the functions of nucleus				
6	Course	After the successful completion of this course students will be able to:				
	Outcomes	CO1: Determine different types of cell membrane and their function like translocation of biomolecules thru' membrane.				
		CO2: Determine the types of organelles and their specific function				
		CO3: Analyse the metabolic activity of the cell and protein transdport process.				
		CO4: Explanation and analysis of bioenergetics and metabolic process				
		CO5: Characterize the functions of Nucleus and its activities thru'				
		cellular organelles				
		CO6: Explanation of the structure and function of cell organelles				
7	Course	Molecular cell biology is a unifying discipline that describes the				
	Description	structure and function of cells in all their genetic, biochemical,				
		developmental, physiological and pathophysiological aspects.				
0	O(1) 11 1					
8	Outline syllabus					



Molecular Composition of Cell Membrane				
Lipid structure and fatty acids, phospholipids forming lipid vesicles,				
membrane proteins, carbohydrate, bacterial outer membrane				
Transport across Cell Membranes ;Ion channels and transport of small				
molecules, channel proteins, carrier proteins; active and passive transport				
of molecues, Antiport				
Endocytosis: Phagocytosis, Receptor mediated Endocytosis				
ER & Protein Sorting				
Endoplasmic Reticulum; targeting protein to ER; Overview of protein sorting; Isolation of rough ER				
Protein folding and processing in ER				
Lysosomes				
Protein Transport				
GPI anchors				
Golgi Apparatus, structure & function				
Protein sorting and export from Golgi, Vesicular transport				
Bioenergetics and Metabolism Metabolism in the matrix of Mitochondria: organization and function;				
Import of mitochondrial matrix protein				
Chloroplast &plastids protein import into chloroplast stroma; import of				
proteins into thyllakoid membrane of chloroplasts;; Electron flow				
through photo system I and II				
Peroxisomes functions				
Internal organization of Nucleus				
Structure of the nuclear envelop; Nuclear Pore complex				
Protein transport to and from Nucleus; functional domain within the				
nucleus				
Cajal bodies ;Nucleolus				
Theory				
Theory				
CA MTE ETE				
30% 20% 50%				
Gerald K., "Cell and Molecular Biology", John Wiley and Sons, 2006.				
1. Cooper G.M., "The Cell: A Molecular Approach", Sinaner Associates,				
2004.				
2. Verma P.S. and Agarwal, V.K., "Cell Biology, Genetics, Molecular				
Biology Evolution and Ecology", S. Chand and Company, 2004.				



BTY613Biological database and their management

Sc	hool: SET	Batch: 2018-2020
	ogram:	Current Academic Year: 2018-19
M.	Tech	
	anch:	Semester: 02
	otechnology	
1	Course Code	BTY613
2	Course Title	Biological database and their management
3	Credits	3
4	Contact	300
	Hours	
	(LTP)	
	Course	Compulsory /Elective/Open Elective
	Status	
5	Course	1. This course surveys a wide range of biological databases and their
	Objective	access tools and enables students to develop proficiency in their use.
		2. The course also focuses on the design of biological databases and
		examines issues related to heterogeneity, interoperability, complex
_		data structures, object orientation and tool integration.
6	Course	
	Outcomes	CO1. Deview different high-girel detahases and welchesed macromaning
		CO1: Review different biological databases and webbased programming
tools to make biological databases accessible.		CO2: Develop databases that store biological information (genome sequence
		database, protein 3D structure database, gene expression profile database,
		molecular interaction database, etc).
		CO3: Develop computing tools for analyzing various kinds of biological and
		experimental data, data mining from databases, computer simulation of
		living systems and so on.
		CO4: Develop ontologies necessary for data and knowledge description of
		databases storing biological functions and integration of the basic databases.
		CO5: Retrieve and interpret the data from different databanks (nucleotide,
		cDNA, rRNA, protein sequence, signal peptide and AIDS virus databanks).
		CO6: Normalize database design and perform experiments using SQL for
		specifying, authorization, viewing, encryption, structure indexing and
		hashing. Design and distribute query processing recovery and operate
		multidatabase and parallel databases systems.
7	Course	To understand how the database is created and the ways to manage it.

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	D	E1	- 1-4-1	aliah aantaina daa historiaal data Tooloo daan daa				
	Description			which contains the biological data. It also clears the				
	0 11 11 1		database design issues and also makes understand the way to protect data					
8	Outline syllabi							
	Unit 1		n to Database					
	A		,	dels, Basic concept of databases, Data				
			independence					
	В		DML, DCL, DDL and structure of database management system					
	C	•		m: Basic and advance concept, Application of ER				
		diagram in d	esigning data	base system				
	Unit 2	Biological D	atabasesI					
	A	Nucleic acid	sequence dat	a banks, Genbank, EMBL, DDBJ				
	В	GenPept, nuc	cleiotide sequ	ence databank				
	C	cDNA datab	ank					
	Unit 3	Biological D	atabasesII					
	A	AIDS virus s	AIDS virus sequence data bank					
	В	rRNA data b	ank					
	С	Protein seque	ence data ban	ks, Signal peptide data bank, NBRFPIR,				
		SWISSPRO	Γ					
	Unit 4	Database Do	Database Design Issues					
	A	Normalizatio	Normalization INF, 2NF, 3NF, 4NF, BCNF and 5NF					
	В	Database des	Database design problems, Security and integrity					
	С	Use of SQL for specifying, authorization, view, encryption, Storage structure						
		indexing and hashing, Different types of file organization						
	Unit 5	Distributed	Database Str	ructure				
	A	Design, trans	sparency and	autonomy, Distributed query processing recovery				
	В			k handling, Multidatabase system				
	С	Parallel data	base concept	and related issues, Web interface to database				
	Mode of		/Practical/Viv					
	examination							
	Weightage	CA	MTE	ETE				
	Distribution	30%	20%	50%				
	Text book/s*	Cohn R. and	Russell J., "I	Biological Databases", VSD Publications, 2012.				
	Other			nardi S., "Biological Data Mining", Chapman and				
	References	Hall, 2009.						
		2. Chen J. and Sidhu A.S., "Biological Database Modeling", Artech						
		House, 2007.						
	l	110400, 2007.						



Elective1

BTY604 Advances in Bioprocess Engineering

Scl	nool: SET	Batch: 2018-20				
Pro	ogram: M.	Current Academic Year: 2018-19				
Te	ch					
Bra	anch: BT	Semester: I (Odd semester)				
1	Course Code	BTY604				
2	Course Title	Advances in Bioprocess Engineering				
3	Credits	3				
4	Contact	300				
	Hours					
	(LTP)					
	Course Status	Compulsory/ Elective /Open Elective				
5	Course Objective	1. To enable students bridge the gap between theoretical concepts and practical aspects in industrial settings				
		 Indepth knowledge and handson laboratory/industrial skills required for employment or for creation of employment in bioprocess engineering. To enable students about nutritional values and increase yield of products by modifying microorganisms. Knowledge to produce antibiotics, vitamins, vaccines and organic solvents using a bioreactor. 				
6	Course Outcomes	 After successful completion of the course students will be able to CO1: Apply mathematical models for calculating substrate uptake, product formation and cell kinetics. CO2: Design strategies for using bioreactors to address different needs of the industry and to conduct scaleup methods for designing bioreactors CO3: Apply the models and mathematical equations to study about the working principles of Bioreactor. 				



7	Course Description	CO4: Understand and apply different strategies for the downstream processing to biomolecules at industrial level. CO5: Understand the industrial production of antibiotics, vitamins, and vaccines. CO6: Understand and apply different bioprocess engineering methods and models for the production and optimization of important microbial products. The course concentrates on bioprocess engineering and bioreactor operation. A considerable part is devoted to the growth analysis using process analytical technology and the evaluation of process data in connection to the generally used cultivation principles.					
8	Outline syllabu						
	Unit 1	Microbial Growth					
	A	Unstructured and structured models for reactor process					
	В	Mathematical models for substrate uptake and product formation					
	С	Kinetics of cell growth, plasmid stability					
	Unit 2	Design of Bioreactors					
	A	Types of microbial and enzyme bioreactors					
	В	Batch, fed batch and continuous processes					
	С	Scaleup of reactor					
	Unit 3	Working of Bioreactor					
	A	Heat transfer and design equations for CSTR fermentor					
	В	Monod model					
	С	Rheology					
	Unit 4	Downstream Processing					
	A	Cell disruption and solvent extraction					
	В	Product recovery					
	С	Sedimentation, floatation, adsorption and chromatography					
	Unit 5	Industrial Applications					
	A	Industrial production of alcohol, citric acid, amino acids, enzymes,					
		antibiotics and steroids					
	В	Microbiology of fermented milk					
	С	Tea, coffee and vinegar fermentation					
	Mode of	Theory/Jury/Practical/Viva					
	examination						
	Weightage	CA MTE ETE					
	Distribution	30% 20% 50%					
	Text book/s*	Doran P.M., "Bioprocess Engineering Principles" Academic Press, 2012.					
	Other References	1. Shuler M.L., "Bioprocess Engineering: Basic Concepts", Pearson Education, 2012.					



	2.	Najafpour	G.D.,	"Biochemical	Engineering	and	Biotechnology",
		Elsevier, 20	007.				

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BTY603Applied Genetic Engineering

Sch	nool: SET	Batch: 2018-2020					
Pro	gram:	Current Academic Year: 2018-19					
M .7	Гесh						
Bra	inch:	Semester: 01					
Bio	technology						
1	Course Code	BTY603					
2	Course Title	Applied Genetic Engineering					
3	Credits	3					
4	Contact	300					
	Hours						
	(LTP)						
	Course Status	Compulsory/Elective/Open Elective					
5	Course	1. To acquire knowledge of principle and techniques involved in genetic					
	Objective	engineering.					
		2. To comprehend the basic strategies of cloning and expression so that may					
		use it for changing the constitution of an organism for human benefit.					
		3. To know about applications of genetic engineering in industry and health					
		sector					
6	Course	CO1: Know and apply the molecular tools, vectors, hosts for genetic					
	Outcomes	manipulation					
		CO2: Comprehend the basic principle of cloning and rDNA technology.					
		CO3: Learn the optimization and technique of DNA amplification by PCR					
		CO4: Analyze gene and protein expression patterns					
		CO5: Create transgenic organisms with desired characteristics using genetic					
		engineering					
		CO6: Understand the basic methods of creating recombinant genes,					
		amplifying the same, creating libraries, engineering proteins and finally					
		apply the knowledge in creating transgenic products with gene delivery					
_		tools					
7	Course	The course covers fundamentals of genetic engineering that leads to specific					
	Description	advanced applications for the benefit of mankind					
8	Outline syllabu						
}	Unit 1	Tools of Genetic Engineering					
	A	Genetic engineering and molecular tools					
	В	Enzymes involved in manipulation of genetic material					
	С	Vectors and host for cloning and cloning process					



Unit 2	Cloning						
A	Cloning and C	Construction of	recombinant DNA				
В	Cloning intera	Cloning interacting genes					
С	Library construction and screening						
Unit 3	In vitro Ampl	lification of D	NA				
A	Polymerase ch	nain reaction ar	nd its types				
В	Cloning of ge	nes by PCR					
С	Optimization	of PCR					
Unit 4	Expression						
A	Expression str	ategies					
В	Vector and ho	Vector and host engineering					
С	Protein engine	eering and gene	e tagging				
Unit 5 Applications							
A	Strategies of gene delivery						
В	Methods for gene expression analysis						
C	Transgenic or	ganisms					
Mode of	Theory/Quiz						
examination							
Weightage	CA	MTE	ETE				
Distribution	30%	20%	50%				
Text book/s*	Brown T.A, '	"Gene Cloning	g and DNA Analysis: An Introduction", John				
	Wiley & Sons, 2010						
Other	1. Old R.W	and Primros	e S.B., "Principles of Gene Manipulation",				
References		Scientific Pub	*				
	-		M. and Plant N., "From Genes to Genomes:				
	Concepts	and Application	ns of DNA Technology", John Wiley, 2011.				



BTY602 Enzyme Technology

Sch	ool: SET	Batch: 2018-2020						
	gram:	Current Academic Year: 2018-19						
M.Tech								
Bra	nch:	Semester: Odd (1 st)						
Bio	technology							
1	Course Code	BTY 602						
2	Course Title	Enzyme Technology						
3	Credits	3						
4	Contact	300						
	Hours							
	(LTP)							
	Course Status	Compulsory / Elective / Open Elective						
5	Course	With this Course the students						
	Objective	1. Will acquire knowledge fundamental Knowledge of Enzymes						
		2. Will get useful exploitation of enzymes physical and kinetic						
		properties						
		3. Use Enzymes biocatalysts in the biotransformations						
		4. Know the Industrial, Research and Therapeutic applications of						
		Enzymes.						
6	Course	After successfully completion of this course students will be able to:						
0	Outcomes	CO1: Basics of Enzymes and its Classification						
	Outcomes	CO2: Evaluate the role of substrates and cofactors in enzyme						
		kinetics.						
		CO3: Predict type of enzyme inhibition by using Lineweaver Burk						
		plot method.						
		CO4: Optimize enzyme catalyzed reactions and compare rate of						
		reactions of enzyme catalyzed and noncatalyzed reactions.						
		CO5: Perform and analyze enzymatic assays using						
		spectrophotometer and microtiter plate reader.						
		CO6: Purify proteins by precipitation and determine protein-protein						
		interaction by coimmunoprecipitation.						
		CO7: Purify native enzymes and compare catalytic activity with						
		engineered enzymes.						
		CO8: Implement the use of enzymes for industrial applications.						
7	Course	This course covers fundamentals to applications necessary for the useful						
	Description	exploitation of enzymes both as tools for the enzymatic analyses and as						



		biocatalysts in the biotransformations on the unique structuralfunctional					
		•	properties of enzymes and its industrial and research utilization.				
8	Outline syllabu						
	Unit 1	Enzymes					
	A		Classification of enzymes				
	В	Properties of					
	С	Factors affe	ecting enzym	natic activity			
	Unit 2	Kinetics of	Kinetics of Enzyme Catalyzed Reaction				
	A	Enzymesub	strate compl	lex			
	В	Enzyme inh	nibition				
	С			ion of enzyme activity			
	Unit 3	Mechanisn	n of Enzyme	ecatalyzed Reaction			
	A		of enzyme				
B Coenzymes and cofactors				ors			
	С	_	on of enzyme				
	Unit 4	Immobilization of Enzymes					
	A	Principle ar	nd kinetics o	f enzyme immobilization			
	В	Multienzyn					
	С			lization and regeneration of cofactors			
	Unit 5		Uses of Enz				
	A			es of enzymes			
	В			eering on enzyme production			
	C	Engineered	enzymes				
	Mode of	Theory					
	examination						
	Weightage	CA	MTE	ETE			
	Distribution	30%	20%	50%			
	Text book/s*			, "Enzymes: Biochemistry, Biotechnology, Clinical			
	0.1	Chemistry"	, Woodhead	Publishing, 2007.			
	Other	1. Copela	and R. A., "I	Enzymes: A Practical Introduction to Structure,			
	References	Mechanism, and Data Analysis", Wiley, 2006.					
		2. Gui	sán J. M., "I	mmobilization of Enzymes and Cells (Methods in			
				', Humana Press, 2010.			



BTY631Molecular Signaling

School:	SET	Batch: 2018-2020
	n: M.Tech	Current Academic Year: 2018-19
	: Biotechnology	Semester: 02
1	Course Code	BTY631
2	Course Title	Molecular Signaling
3	Credits	4
4	Contact Hours (LTP)	310
	Course Status	Compulsory /Elective/Open Elective
5	Course Objective	 To understand how communication takes place between different cells in the body. To elucidate the signal transduction pathways involved in several diseases which is important to define the new target for drug development.
6	Course Outcomes	CO1: Determine the types of communication between cells and correlate deregulation of extracellular matrix with occurrence of different diseases. CO2: Analyse the progression of signals inside the cell by identify the role of secondary messengers in signalling pathways. CO3: Perform covalent modification (phosphorylation) by using serine/threonine and tyrosine protein kinases thus understand pathways in cells during different types of stress/ signalling. CO4: Understand the neuronal signalling in correlation with its regulatory pathways. CO5: Demonstrate the role played by tumour suppressor genes and oncogenes thus recognize the roles played by proapoptotic, antiapoptotic proteins and caspases in apoptosis. CO6: To identify the possibilities, efficacy and potency of therapeutic drugs in cell signalling pathways for disease treatments.
7	Course Description	To understand how communication takes place between different cells in the body. To elucidate the signal transduction pathways involved in several diseases which is important to define the new
0	Outling avillate	target for drug development.
8	Outline syllab	us



Unit 1	Cellular Co	mmunication	1					
A	Introduction	to cell signal	ing.					
В	Intercellular	Intercellular communication and its types						
С	Extracellula	r matrix, Neur	otransmitter	s and Neu	ırohormone	es		
Unit 2	Signal Tran	sduction						
A	Receptors an	nd its types.						
В	Gprotein co	upled receptor	mediated sig	gnalling.				
C	Modulation	of different si	gnalling by s	econdary	messenger	rs.		
Unit 3	Protein Kin	Protein Kinases and their pathways						
A		s and inhibitor		-			of	
В	Protein Kina	ise A pathway	and Regula	tion of PI	3K/Akt pat	hway.		
С	MAPK casc	ades.						
Unit 4	Siganling in	Siganling in Plants						
A		Phytohormones and signaling mechanisms						
В	Phytochrom	es and Crypto	chrome					
C	·	ention in plant	s.					
Unit 5	Signalling in	n Cancer						
A	Oncogenes a	and tumour su	ppressor ger	ies.				
В	1 0	ression and m						
C		nd therapeutic	intervention	for treati	ing cancer.			
Mode of examina		//Practical/Viv	'a					
Weighta	ge CA	MTE	ETE					
Distribut	ion 30%	20%	50%					
Text boo		Krauss G., "Biochemistry of Signal Transduction and Regulation", WileyVCH, 2008.						
Other Reference		1. Hancock J.T., "Cell Signalling", Oxford University Press, 2010.						
		perts B.D., Ki sduction", Ac			n P.E.R., "	Signal		



BTY630 Cell and Tissue Engineering

Scł	nool: SET	Batch: 2018-2020					
Pro	ogram:	Current Academic Year: 2018-19					
Μ.	Tech						
Bio	technology	Semester: Even(2 nd)					
1	Course Code	BTY630					
2	Course Title	Cell and Tissue Engineering					
3	Credits	4					
4	Contact	310					
	Hours						
	(LTP)						
	Course Status	Compulsory /Elective/Open Elective					
5	Course	1. To Study cell, tissue culture, media component					
	Objective	2. To Study Cell Viability and Kinetics					
		3. To Study Cell cloning, cell genetics					
		4. To Study industrial medical and agricultural applications of cell and					
		tissue engineering.					
6	Course Outcomes	After successfully completion of this course students will be able to:					
		CO1: Understand basics of Cell and Tissue culture, evaluate media and aseptic techniques of establishing primary and Secondary cell cultures. CO2: Understand the concepts and Mechanism of Cell Viability adherence, calculate growth kinetics parameters and apply cryopreservation technique for long term storing of cells. CO3: Evaluate Cell Characteristics, Cell Signaling, genetics, establish a continuous cell line from cells of different origin and determine their nutrient and environment requirements CO4: Understanding Cell Cloning for Tissue Engineering and Stem Cell Therarpy, Biomaterials for Cells CO5: Understand Applications of Cell and Tissue Engineering for Industrial, Agriculture medical applications CO6: Acquiring Aquaintence of Cell Culture Technology by studying cell, tissue culture, media component, cloning, cell genetics and large scale industrial, agriculture and medical applications of cell and tissue engineering.					
7	Course	To acquire a fundamental and advanced knowledge of Cell and Tissue					
	Description	Culture Technology by studying cell, tissue culture, media component, cloning, cell genetics and large scale industrial, agriculture and medical					



		applications of cell and tissue engineering.					
8	Outline syllabu						
	Unit 1			nd Tissue Culture			
	A	History of 0	Cell Culture,	Cell, Tissue and organ culture, Culture procedures			
	В	Culture me	dia and grow	th conditions, primary and Secondary cultures			
	С	Establishme laboratory a		ntenance of cell lines and Risks in a tissue culture			
	Unit 2	•	cs and Viab	sility			
	A			, Characterization of cultured cells morphology of			
	В	cell adhesic	_	ion, differentiation, Kinetics involved in growth of			
	С	Cell viabili	ty, Methods	for testing cell viability, Cytotoxicity assays			
	Unit 3		and Cell C				
	A	Introduction	n to Stem Ce	ells and its Types			
	В	Methods of	Cloning of	Stem Cells			
	C	Stem Cells	Application	ls .			
	Unit 4	Biomateria	ls for Tissu	e Engineering			
	A		-	s Of Biomaterials ,Surface, Bulk, Mechanical And			
	D	Biological		D' 1 ' 1 A 1 G (1 (' M (' 1 D') 1			
	B C			Biological And Synthetic Materials, Biopolymers			
	C	Application		' OCD' ('I D I OCN (I I			
	TT *4 =			ions Of Biomaterials, Role Of Nanotechnology.			
	Unit 5			nd Tissue Engineering			
	A			of Cell and Tissue Engineering			
	В			cations of Cell and Tissue Engineering			
	С		griculture Ir	ndustrial applications of Cell and Tissue Engineering			
	Mode of	Theory					
	examination						
	Weightage	CA	MTE	ETE			
	Distribution	30%	20%	50%			
	Text book/s*	Butler M., "Animal Cell Culture and Technology", Garland Science, 2008. Bhojwani S.S., Dantu P.K., "Plant Tissue Culture: An Introductory Text", Springer, 2013.					
	Other	Jenkins N.,	"Animal Ce	ll Biotechnology: Methods and Protocols", Humana			
	References	Press, 2006. Freshney I.R., "Culture of Animal Cells: A Manual of Basic Technique", Wiley, 2005.					



BTY606 Applied Bioinformatics

Sc	hool: SET	Batch: 2018-2020							
	ogram:	Current Academic Year: 2018-19							
	Tech								
	anch:	Semester: 02							
	otechnology								
1	Course Code	BTY606							
2	Course Title	Applied bioinformatics							
3	Credits	3							
4	Contact	300							
	Hours								
	(LTP)								
	Course	Compulsory / Elective / Open Elective							
	Status								
5	Course	1. To acquire an advanced knowledge of bioinformatics tools used for							
	Objective	designing and analyzing in silico experiments and different techniques.							
		2. To attain knowledge about data storage model, retrieval of information							
		and integration. To learn the procedure of sequence alignment and							
		phylogenetic analysis by using different online and offline tool along							
		with their algorithms.							
		3. To understand about gene organization, genome sequencing, gene							
		prediction methods and motif search methods. To have a clear cut idea							
		about bioinformatics scope, concepts and major databases/tools/softwares							
		with their algorithms used for various application							
6	Course	CO1: Analyze sequence similarity search using BLAST.							
	Outcomes	CO2: Examine phyolgenetic relationship using clustal and parsimony.							
		CO3: Assess motif consensus by Markov model.							
		CO4: Identify regulatory sequence by Meme.							
		CO5: Determine structure of biomolecules by software (Pymol, Rasmol) and							
		database.							
		CO6: Compute structure of biomolecules using modeling and docking.							
		Perform microarray and protein array analysis for drug target identification							
7	Course	and gene prediction. To acquire a fundamental knowledge of basic computational biology by							
/	Description	studying, designing and analyzing <i>insilico</i> experiments. To learn the							
	Describtion	studying, designing and analyzing msilico experiments. To learn the							

Prepared by Department of Biotechnology



	procedure of sequence alignment and its application in molecular								
				and different techniques used for gene prediction					
			of biological						
8	Outline syllabi								
	Unit 1	Sequencealignment Related Problems							
	A	•	Sequence databases, Similarity matrices, pairwise alignment, BLAST						
	В		Sequence assembly, multiple sequence alignment						
	С		Clustal, phylogenetics: distance based approaches, parsimony						
	Unit 2	Pattern Analysis in Sequences							
	A	Motif representation: consensus, regular expressions, Markov model							
	В			ification using Meme					
	С		_	based finding, sequence motifbased finding					
	Unit 3		ated Problen						
	A	Representation	on of molecul	ar structures (DNA, mRNA, protein), secondary					
		structures, do	omains and mo	otifs					
	В	Visualization	software (Py	mol, Rasmol)					
	С	Experimenta	l determinatio	n of structures (Xray crystallography, NMR),					
		Structure dat	abases						
	Unit 4	Structurerelated ProblemsII							
	A	Ab initio stru	icture predicti	on: force fields, backbone conformer generation					
		by Monte Ca	rlo approache	S					
	В	Protein struc	ture prediction	by comparative modeling approaches (homology					
		modelling, th							
	C			mputeraided drug design (pharmacophore					
		identification	ı), QSAR						
	Unit 5	Systemwide							
	A	Transcripton							
	В			pression profiles, data analysis, SAGE					
	C	•		ics: ¹³ C NMR based metabolic flux analysis					
	Mode of	Theory/Jury	/Practical/Viv	a					
	examination								
	Weightage	CA	MTE	ETE					
	Distribution	30%	20%	50%					
	Text book/s*	Jin X., "Esse	ential Bioinfo	rmatics", Cambridge University Press, 2006.					
	Other	1. Mount D.	W., "Bioinfor	matics: Sequence and Genome Analysis", Cold					
	References			ory Press, 2004.					
		2. Baxevanis	A., Ouellette	F.B.F., "Bioinformatics: A practical guide to					
		the analy	the analysis of genes and proteins", WileyInterscience, 2004.						
		3. Bourne P.	E., Gu J., "St ı	ructural Bioinformatics", WileyBlackwell, 2009.					



BTY607 Immunotechnology

School: SET		Batch: 2018-2020			
Program:		Current Academic Year: 2018-19			
M.Tech.					
Br	anch:	Semester: 02			
Bi	otechnology				
1	Course	BTY607			
	Code				
2	Course	Immunotechnology			
	Title				
3	Credits	3			
4	Contact	300			
	Hours				
	(LTP)				
	Course	Compulsory / Elective /Open Elective			
	Status				
5	Course	1. Understand anatomy of immune system, immunity and molecular basis			
	Objective	of various immune responses.			
		2. Discuss about the structure and function of antibody and MHC.			
		3. Understand and discuss the various immunotechniques, immunization			
		and vaccines.			
6	Course	CO1: Describe immune system, immunity and immune responses			
	Outcomes	CO2: Explain structure and function of antibodies, BCR, TCR and MHC;			
		AgAb reaction			
		CO3: Discuss about the molecular basis of immune response.			
		CO4: Explain various techniques in immunology.			
		CO5: Demonstrate the principle behind the immnunization; vaccine and its			
		types.			
		CO6: Explain the organization and functioning immune system, immunity,			
		vaccine, vaccination and immunological techniques.			
7	Course	The course will help students to acquire a fundamental working knowledge of			
	Description	the basic principles of immunology; to begin to understand how these			
		principles apply to the process of immune function; and to develop the ability			
		to solve problems in clinical immunology by making use of existing tools and			
		techniques			
8	8 Outline syllabus				



Unit 1	Anatomy of Immune System			
A	Cellmediated and humoral immunity; Innate and acquired immunity			
В	Complement and inflammatory responses			
С	of primary and secondary lymphoid organs			
Unit 2	Antibody and MHC			
A	Structure and function of immunoglobulins			
В	Major histocompatability complex and Complement system			
С	BCR, TCR and antigenantibody reaction			
Unit 3	Molecular 1	Basis of Imm	une Response	
A	Activation of	of Tlymphocy	tes and Blymphocytes	
В	Cellmediated, antibodymediated and macrophagemediated cytotoxicity			
С	Cytokine release and their role in immune regulation			
Unit 4	Techniques in Immunology			
A	RIA and types of ELISA			
В	Immunofluorescence and immunoelectron microscopy			
С	CMI Techniques			
Unit 5	Vaccinology			
A	Vaccination and types of vaccines			
В	Recombinant DNA and protein based vaccines, peptide and conjugate			
	vaccines			
С	Antibody engineering, catalytic antibody and generation of immunoglobulin			
	gene libraries			
Mode of Theory/Jury/Practical/Viva			va	
examination		T		
Weightage	CA	MTE	ETE	
Distribution	30%	20%	50%	
Text book/s* Kindt T.J., Osborne B.A. and Goldsby R.A. (2006) Kuby Imm Freeman Other 1. Delves P.J, Martin S.J., Burton D.R. and Roitt I.M., (20			and Goldsby R.A. (2006) Kuby Immunology, W.	
References	Essential Immunology, Wiley			
	2. Paul B.W.E, "Fundamental Immunology", Lippincott Williams and			
	Wilk	ins, 2008.		



BTY 632 Computer Aided Drug Design

<u> D1</u>	BTY 632 Computer Aided Drug Design				
School: SET		Batch: 2018-20			
Program: M Tech		Current Academic Year: 2018-19			
Branch: Genetic		Semester: II			
Eng	gineering				
1	Course Code	BTY 632			
2	Course Title	Computer Aided Drug Design			
3	Credits	3			
4	Contact Hours	3-0-0			
	(L-T-P)				
	Course Status	DE			
5	Course	Upon completion of this syllabus, the student can able to understand			
	Objective	1. Role of Bioinformatics/Chemo-informatics in drug designing and			
		discovery process.			
		2. Different CADD techniques and their importance and applications.			
		3. Various strategies to design and develop the drug-like/lead-like molecules.			
6	Course	CO1: To understand the basics of bioinformatics, chemo-informatics and			
Outcomes		how useful for drug designing and discovery process.			
		CO2: Acquire the knowledge about protein structure prediction methods,			
		structure visualizations and their importance.			
		CO3: Understand the principle, types and various applications of			
		computer aided drug designing and discovery process.			
		CO4: Explore the concept and SAR, QSAR and their importance in ligand			
		optimization.			
		CO5: Understand the principle and applications of molecular dynamics simulation.			
		CO6: Overall understanding the concept and applications for computer aided drug designing and discovery process.			
7	Course	This syllabus covers the various components of computer aided drug			
'	Description	designing and discovery process namely protein structure preparation,			
	Description	ligand structure preparation, structural databases, virtual screening			
		techniques, SAR/QSAR, molecular mechanics and molecular dynamics			
		simulation.			
8					
	Unit 1 Introduction				
		History of drug design, Stages of drug discovery and development; Drug			
	properties, likeness; Role of Bioinformatics and Chemo-informatics				
	B Classification of Protein Structures – Primary, Secondary,				
		secondary, Tertiary and Quaternary; Active Sites; Allosteric Sites;			
	Domains; Fold; Motif;				
	С	Structural databases- PDB, PDBSUM, SCOP, CATH; Chemical and Drug			
	-	Molecule Databases – PubChem, Zinc and DrugBank			
	I	,			



	Unit 2	Preparation of Protein Structure		
	A	Introduction to <i>in silico</i> and experimental protein structure determination		
		methods;		
	В	In silico Structure Prediction - Homology Modeling; Threading; Fold		
		Recognition. Ab initio modeling;		
	С	Model refinement and validation; Prediction of Binding site; Structure		
		Visualization and Analysis tools.		
	Unit 3	High throughput Virtual Screening and Molecular Docking		
	A	Types of Virtual Screening methods; Structure Based Virtual Screening;		
		Ligand Based Virtual Screening		
	В	Library design; Concept of pharmacophore mapping and pharmacophore based Screening;		
	С	Molecular Docking: Rigid and Flexible docking; Analysis of Protein-		
		Ligand interactions.		
	Unit 4	Quantitative Structure Activity Relationship (QSAR)		
	A SAR versus QSAR, History and development of QSAR,			
		physicochemical parameters,		
	В	experimental and theoretical approaches for the determination of		
		physicochemical parameters such as Partition coefficient, Hammet's		
		substituent constant and Tafts steric constant.		
	C	Hansch analysis, Free Wilson analysis, 3D-QSAR approaches like		
		COMFA and COMSIA.		
	Unit 5	Molecular Mechanics and Molecular Dynamics Simulations		
	A	General features of molecular mechanics; Energy Minimization - loc		
		and global energy minima, saddle point, applications.		
	В	Molecular dynamics simulation		
	С	Understanding the structural stability of protein and protein-ligand complex.		
	Mode of	Theory		
	examination			
	Weightage	CA MTE ETE		
	Distribution	30% 20% 50%		
	Text book/s*	Lednicer, D. (1998) "Strategies for Organic Drug Discovery Synthesis		
		and Design"; Wiley International Publishers.		
	Andrew R. Leach (2001). Molecular Modeling – Principles and			
	References	Applications. Second Edition, Prentice Hall, USA		



BTY 633 Animal Cell Technology

		Cell Technology			
School: SET		Batch: 2018-20			
Program: M Tech		Current Academic Year: 2018-19			
Branch: Genetic		Semester: II			
Eng	gineering				
1	Course Code	BTY 633			
2	Course Title	Animal cell Technology			
3	Credits	3			
4	Contact Hours	3-0-0			
	(L-T-P)				
	Course Status	DE			
5 Course This course will result in understanding of					
	Objective	1. Students will understand gene transfer technologies for animals and animal			
		cell lines			
		2. To impart the knowledge on basic tissue culture techniques;			
		3. To apply the state of art knowledge of subject for the production of			
		transgenic animals and production modern drug delivery or vaccination			
-	methods.				
6 Course		After successfully completion of this course students will be able to:			
	Outcomes	CO1: Demonstrate foundational knowledge of Cell culture techniques and competence in laboratory techniques			
		CO2: Understand various types of media and supplements required for			
		animal cell culture.			
		CO3: Familiarize with basic concept of cell lines, immobilization and			
		maintenance of cell culture.			
		CO4: Understand basic concept of scale up of animal cell culture.			
		CO5: Acquire knowledge in tissue engineering and its applications by			
		various methods			
		CO6: Acquire adequate knowledge in the animal cell culture, genetically			
		modified organisms and their beneficial uses			
		This course provides a brief understanding about the animal cell			
	Description	techniques, their set up requirements, scale up and their applications in			
	_	various fields.			
8 Outline syllabus					
Unit 1 Animal Cell Culture		Animal Cell Culture			
	A	Introduction, importance, history of cell culture development, different tissue			
		culture techniques including primary and secondary culture, continuous cell			
		lines, suspension culture.			
B		Advantages and limitations of animal cell culture, genetic engineering of animal			
C		cells and their applications.			
	C	Risks in a tissue culture laboratory and safety and biohazards.			



	Unit 2	Animal Cell Culture Requirements			
	A	Facilities for animal cell culture, infrastructure, equipment, culture vessels.			
	В	Different types of cell culture media, growth supplements, serum free media,			
				ll culture reagents	
	С	Biology and c	haracterization	of cultured cells, cell adhesion, proliferation,	
		differentiation	, morphology	of cells and identification.	
	Unit 3	Primary cell culture techniques			
	A	Mechanical disaggregation, enzymatic disaggregation, separation			
		viable and nor	n-viable cells.	Mass culture of cells, manipulation of cell line	
		selection, type	es of cell lines,	maintenance of cell lines	
	В	immobilizatio	n of cells ar	nd its application, synchronization of cell	
				n, production of secondary metabolites,	
		biotransformation,			
	C Induction of cell line mutants and mutations, cryopreserv			s and mutations, cryopreservation, germplasm	
	conservation and establishment of gene banks.				
	Unit 4	Animal Cell Culture Scale-up			
	A	Scale up in suspension, stirrer culture, continuous flow culture, air-lift			
		fermenter culture			
	В	Scale up in monolayer using Roller bottle culture, multi-surface culture,			
		multi-array disks, spirals and tubes			
	С	Monitoring of cell growth and cell death.			
	Unit 5	Tissue engineering and its applications			
	A	Design and engineering of tissues, tissue modeling. Embryonic stem cell			
		engineering, ES cell culture to produce differential cells.			
	В	Human embryonic stem cell research and embryo micromanipulation			
	С	Transgenic animals, and xenotransplantation			
	Mode of	Theory		-	
	examination				
	Weightage	CA	MTE	ETE	
	Distribution	30%	20%	50%	
Text book/s* Freshney I. Culture of Animal Cells: A Manual of Basic Techr				nal Cells: A Manual of Basic Technique, 5th	
	Edition Publisher: Wiley-Liss, 2005 ISBN: 0471453293				
	Other	Nigel Jen, Animal Cell Biotechnology: Methods and protocols, Humana			
	References Press				