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S.P. Mandali's

Ramnarain Ruia Autonomous College

(Affiliated to University of Mumbai)



Syllabus for T.Y.B.Sc Program: BSc (Microbiology) Program Code: RUSMIC

(Credit Based Semester and Grading System for the academic year 2019-2020)

SEMESTER V

COURSE				
CODE	UNIT	TITLE	CREDITS	LEC/WEEP
RUSMIC 501		MICROBIAL GENETICS	2.5	4
	I	BRANCHES OF GENETICS, PLASMIDS AND TRANSPOSONS		
	II	DNA REPLICATION		
	ш	MUTATION AND REPAIR	C	O^{\star}
	IV	HOMOLOGOUS RECOMBINATION &GENETIC	S	
RUSMIC 502			2.5	4
	I	GENETICS OF PATHOGENICITYANDSTUDY OF INFECTIOUS DISEASES-I		
	II	STUDY OF INFECTIOUSDISEASES-II		
	Ш	STUDY OF INFECTIOUS DISEASES-III		
	IV	CHEMOTHERAPY OF INFECTIOUS DISEASES		
RUSMICP		PRACTICALS BASED ON ABOVE TWO COURSES	3	4
501	<u> </u>			

SEMESTER V

COURSE				
CODE	UNIT	TITLE	CREDITS	LEC/WEEK
RUSMIC 503		MICROBIAL BIOCHEMISTRY: PART-I	2.5	4
	I	BIOLOGICAL MEMBRANES & TRANSPORT		
	II	BIOENERGETICS & BIOLUMINESCENCE		
	III	METHODS OF STUDYING METABOLISM & CATABOLISM OF CARBOHYDRATES	S	
	IV	FERMENTATIVE METBOLISM & ANABOLISM OF CARBOHYDRATES	\mathcal{S}	
RUSMIC 504		BIOPROCESS TECHNOLOGY	2.5	4
	I	UPSTREAM PROCESSING		
	II	FERMENTER EQUIPMENT AND CONTROL		
	- 111	DOWNSTREAM PROCESSING		
	IV	BIOINSTRUMENTATION AND IPR		
		PRACTICALS BASED ON ABOVE	3	4
RUSMICP 502		TWO COURSES	Ŭ	

SEMESTER VI

SEMESTER VI

COURSE				
CODE	UNIT	TITLE	CREDITS	LEC/WEE
RUSMIC 603		MICROBIAL BIOCHEMISTRY PART II	2.5	4
	I	LIPID METABOLISM & CATABOLISM OF HYDROCARBONS		
	II	METABOLISM OF PROTEINS AND NUCLEIC ACIDS		
	ш	METABOLIC REGULATION	C	$\mathbf{\mathcal{Y}}$
	IV	PROKARYOTIC PHOTOSYNTHESIS & INORGANIC METABOLISM	S	
RUSMIC 604			2.5	4
	I	INDUSTRIAL FERMENTATIONS-I		
	П	INDUSTRIALFERMENTATIONS-II		
	ш	INDUSTRIALFERMENTATIONS III		
	IV	BIOASSAYS, QUALITY ASSURANCE		
RUSMICP 602		PRACTICALS BASED ON ABOVE TWO COURSES	3	4
	201	JV.		

Course Code: RUSMIC 501 Course Title: MICROBIAL GENETICS Academic year 2019-20

Learning Objectives:

With a background of nucleic acids in FYBSc and Mendelian genetics, DNA structure and transcription, translation and genetic code at the SYBSc level, the undergraduate T.Y. B.Sc. Microbiology course under the Paper on Microbial Genetics introduces the learner to the underlying theories of genetics by elaborating both conceptual and practicaltoolsforquantitative genetics and use of model organisms. It elaborates on extrachromosomal DNA – plasmids and on nature and role of transposons. The course then deals in detail with generating, processing and understanding biological genetic information. It develops knowledge of the underlying theories of genetics by elaborating on various concepts related to DNA replication, mutations and genetic exchange among prokaryotes.

- Understand population and quantitative genetics and get introduced to different model organisms used in genetic studies.
- Understand different natural plasmids and transposons present in prokaryotes
- Understand the molecular mechanism involved in DNA replication
- Understand how to identify and classify mutations in DNA followed by mechanism of DNA repair
- Understand basic concepts of homologous recombination and genetic exchange among prokaryotes

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Course Code	Title	Credits
RUSMIC 501	MICROBIAL GENETICS	2.5 Credits(65 lectures)
Unit I	BRANCHES OF GENETICS, PLASMIDS, TRANSPOSONS	15 lectures
	1.1. Overview of branches of Genetics	4
	i.Transmission, Molecular,	\sim
	ii.Population Genetics: Hardy-Weinberg Law- principle and	\mathbf{O}
	violation of assumptions (Mutation, Migration, Genetic Drift,	
	Natural Selection)	
	iii.Quantitative Genetics: Characteristics, concept of Heritability, QTLs, Response to selection	
	Tieriability, & LS, Response to selection	
	1.2. Model Organisms	
	i.Characteristics of a model organism	3
	ii.Examples of select model organisms used in study: <i>E.coli,</i>	
	Yeast, Mouse.	
	1.3. Plasmids	4
	a. Physical nature	
	 b. Detection and isolation of plasmids c. Plasmid incompatibility and Plasmid curing 	
	 c. Plasmid incompatibility and Plasmid curing d. Cell to cell transfer of plasmids 	
	e. Types of plasmids	
	i.Resistance Plasmids	
	ii.Plasmids encoding Toxins and other Virulence	
	characteristics	
	iii.col factor	
	iv.Degradative plasmids	
	1.4.Transposable Elements in Prokaryotes	4
	a. Insertion sequences	
, D	b. Transposons	
	i.Types	
	ii.Structure and properties	
14.	iii.Mechanism of	
	iv. Transposition	
•	v.Integrons	

Unit II	DNA REPLICATION	15 lectures
	2.1. Historical perspective – conservative, dispersive, semi-conservative, Bidirectional and semi-discontinuous replication	4
	2.2. Prokaryotic DNA replication - Details of molecular mechanism involved in Initiation, Elongation and Termination	4
	2.3. Enzymes and proteins associated with DNA replication- primase, helicase, topoisomerase, SSB, DNA polymerases, ligases, Ter and Tus proteins	4
	2.4. Eukaryotic DNA replication Molecular details of DNA synthesis, replicating the ends of the chromosomes	2
	2.5. Rolling circle mode of replication	1
Unit III	Mutation and Repair	15 lectures
	 3.1. Mutation 3.1. a. Terminology: alleles, homozygous, heterozygous, genotype, phenotype, Somatic mutation, Germline mutation, Gene mutation, Chromosome mutation, phenotypic lag, hotspots and mutator genes 	1
	3.1. b. Fluctuation test.	1
	3.1. c. Types of mutations: Point mutation, reverse mutation, suppressor mutation, frameshift mutation, conditional lethal mutation, base pair substitution, transition, transversion, missense mutation, nonsense mutation, silent mutation, neutral mutation, pleiotropic mutations.	1
SP	3.1.d. Causes of mutation: Natural/spontaneous mutation replication error, depurination, deamination. Induced mutation: principle and mechanism with illustrative diagrams for –	5
N	 i.Chemical mutagens- base analogues, nitrous acid, hydroxyl amine, intercalating agents and alkylating agents. ii.Physical mutagen iii.Biological mutagen (only examples) 2.1 a Amage test 	
	3.1.e. Ames test3.1.f. Detection of mutants	1

	3.2. DNA Repair	
	a. Mismatch repair	5
	b.Light repair	°
	c.Repair of alkylation damage	
	d.Base excision repair	
	e.Nucleotide excision repair	
	f.SOS repair	
Unit IV	Genetic Exchange	15 lectures
	4.1. Gene transfer mechanisms in bacteria &	
	homologous recombination	
	4.1. a. Transformation	4
	i. Introduction and History	
	ii. Types of transformation in prokaryotesNatural	
	transformation in <i>Streptococcus pneumoniae</i> ,	
	Haemophilus influenzae and Bacillus subtilis	
	iv. Problems based on transformation.	-
	4.1. b. Conjugation	5
	i. Discovery of conjugation in bacteria	
	ii. Properties of F plasmid/Sex factor	
	iii. The conjugation machinery	
	iv. Hfr strains, their formation and mechanism of	
	conjugation	
	v. F' factor, origin and behaviour of F' strains,	
	Sexduction.	
	vi.Mapping of bacterial genes using conjugation (Wolman	
	and Jacob experiment).	
	vii.Problems based on conjugation	
	4.1.c. Transduction	3
	i.Introduction and discovery	
	ii.Generalised transduction	
	iii.Use of Generalised transduction for mapping genes	
	iv.Specialised transduction	
	v.Problems based on transduction	
	4.2. Recombination in bacteria	
	4.2.a. General/Homologous recombination	3
	i.Molecular mechanism	
110	ii.Holliday model of recombination	
1/2	b. Site -specific recombination	

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- 2. Benjamin A. Pierce (2008), Genetics a conceptual approach, 3rd ed., W. H. Freeman and company.
- 3. R. H. Tamarin, (2004), Principles of genetics∥, Tata McGraw Hill.
- 4. D,.Nelson and M.Cox, (2005), Lehninger's Principles of biochemistry , 4th ed. Macmillan worth Publishers.
- 5. M.Madigan, J.Martinko, J.Parkar, (2009), Brock Biology of microorganisms 12th ed., Pearson Education International.
- 6. Fairbanks and Anderson, (1999), Genetics∥, Wadsworth Publishing Company.
- 7. Prescott, Harley and Klein, Microbiology, 7th edition McGraw Hill international edition.
- 8. Robert Weaver, Molecular biology, 3rd edn. McGraw Hill international edition.
- 9. Nancy Trun and Janine Trempy, (2004), Fundamental bacterial genetics, Blackwell Publishing
- 10. Snustad, Simmons, Principles of genetics, 3rdedn. John Wiley & sons, Inc.
- 11. Stanier, Ingraham. |General Microbiology|,5 edn.
- 12. Benjamin Lewin, Genes IXI, , Jones and Bartlett publishers.
- 13. JD Watson, Molecular biology of the genel, , 5thedn.

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Course Code: RUSMIC 502

Course Title: MEDICAL MICROBIOLOGY Academic year 2019-20

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Learning objectives:

Classical medical microbiology is the study of aetiology, transmission, pathogenesis, clinical manifestations, laboratory diagnosis, prophylaxis and treatment of various bacterial, viral, fungal and parasitic infections. The course on Medical Microbiology introduces the students to all these parameters of representative diseases from each category. The course also includes one of the most important areas of modern medical microbiology that is -understanding genetic modification and pathogen evolution.

As a part of understanding chemotherapeutic agents for destruction of pathogens, the students are introduced to different classes of chemotherapeutic agents and their mechanisms of action. As development of resistance to antibiotics is a very burning issue in the field of clinical microbiology, the syllabus also includes mechanisms of resistance to drugs.

- Understand modern alternatives to Koch's Postulates and understand Genetic modification and pathogen evolution
- Study pathogenesis and clinical features of different diseases
- Comment on the mode of transmission, epidemiology and therefore modes of prophylaxis of these diseases
- Given a few key clinical features, identify the likely causative agent.
- Comment on the methods of diagnosis of the disease.
- Correlate classes of antibiotics with their mechanism of action
- Comment on drug resistance mechanisms
- Evaluate drugs and antibiotics for their efficacy

Course Code	Title	Credits
RUSMIC502	MEDICAL MICROBIOLOGY	2.5 Credits(65 lectures)
Unit I	GENETICS OF PATHOGENICITYAND STUDY OF INFECTIOUSDISEASES-I	15 lectures
	1.1.Associating Microbes to disease	02
	1.1.1: Koch's Postulate and modern alternatives to it 1.1.2: Molecular Koch's postulates	\sim
	1.1.2. Molecular Roch's postulates	
	1.2: Genetic modification and pathogen evolution:	03
	1.2.1: Point mutations, gene duplication, chromosomal	
	rearrangements, phase variation and antigenic variation	
	1.2.2: Horizontal gene transfer through Mobile genetic elements	
	1.2.3: Pathogenicity islands	02
	1.3: Sample collection, transport and processing	
	and diagnostic cycles	
	1.4. Study of Infectious Diseases-I	
	(with Emphasis on Characteristics of the Aetiological	08
	Agent, Pathogenesis & clinical features, Laboratory	00
	Diagnosis and Prevention)	
	Study of Respiratory diseases	
	1.4.1. Strep throat by S. pyogenes	
	1.4.2. Diphtheria	
	1.4.3. Common cold 1.4.4. Tuberculosis	
0	1.4.5. Pneumonia caused by <i>K</i> .pneumoniae	
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Unit II	STUDY OF INFECTIOUS DISEASES II	15 lectures
	(With emphasis on cultural characteristics of the	
<u>N</u> .	aetiological agent,pathogenesis, laboratory diagnosis	
	and prevention)	05
~	2.1 Study of skin infections	
	2.1.1 Leprosy	
	2.1.2 Fungal infections- Oral Thrush, Dermatophytosis	

2.1.3 Pyogenic skin infections caused by	
Pseudomonas, S.pyogenes and S. aureus.	
	08
2.2 Study of gastrointestinal tract infections	
2.2.1 Enteric fever- Salmonella	
2.2.2 Shigellosis	
2.2.3 Rotavirus diarrhoea	
2.2.4 Dysentery due to Entamoeba histolytica	
2.2.5 Infections due to pathogenic <i>E.coli</i> strains	02
2.3 Study of urinary tract infections	
Predisposing factors, List of causative agents,	\sim
Pathogenesis and laboratory diagnosis	
Unit III STUDY OF INFECTIOUS DISEASES III	15 lectures
(With emphasis on cultural characteristics of the	
aetiological agent, pathogenesis, laboratory diagnosis	
and prevention)	
3.1 Study of vector-borne infections- Rickettsial	03
diseases (Tabular form), Malaria	
3.2 Study of sexually transmitted infectious	07
diseases	
a. Syphilis	
c. Gonorrhoea	05
3.3 Study of central nervous system infectious	
diseases	
a. Tetanus	
b. Polio	
a. Meningococcal meningitis	
Unit IV CHEMOTHERAPY OF INFECTIOUS AGENTS	15 lectures
Attributes of an ideal chemotherapeutic agent and	03
related definitions	
ction and testing of antibiotics for bacterial isolates by	
Kirby-Bauer method and other assays (E-test &	
Checker Board Assay)	
4.2: Mode of action of antibiotics on-	08
a. Cell wall (Beta-lactams- Penicillin and	
a. Cell wall (Beta-lactams- Penicillin and Cephalosporins, Carbapenems)	
a. Cell wall (Beta-lactams- Penicillin and	

d. Tetracycline and Chloramphenicol) e. Nucleic acid (Quinolones, Nalidixic acid, Rifamycin)	
f. Enzyme inhibitors (Sulfa drugs, Trimethoprim)	01
4.3: List of common antibiotics used for treating viral, fungal and parasitic diseases, New antibiotics	03
4.4: Mechanisms of drug resistance- Its evolution, pathways and origin	

- 1. Jawetz, Melnick and Adelberg's Medical Microbiology, 26th Edition, Lange publication
- 2. Bacterial Pathogenesis A molecular approach Abigail Salyer And Dixie Whitt 2nd Ed ASM press
- 3. Ananthanarayan and Panicker's, Textbook of Microbiology, 9th edition
- 4. Goering, Dockerel et al, Mim's Medical microbiology, 5th Ed 2013, Saunders
- 5. Baron Samuel , Medical Microbiology, 4th edition
- 6. http://www.ncbi.nlm.nih.gov/books/NBK7627/

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Course Code: RUSMIC503 Course Title: MICROBIAL BIOCHEMISTRY PART I Academic year 2019-20

Learning objectives:

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This course is designed for T.Y.B.Sc. Microbiology students such that the students achieve a basic understanding of solute transport and metabolism. The course has been designed to expose students to methods of studying energy generation, fermentative metabolism as well as anabolism.

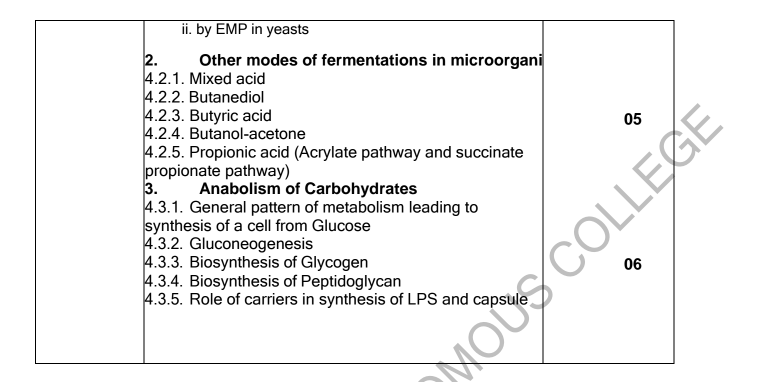
There has been a lot of importance attached to biochemical reactions in living cells. The student must be exposed to the mechanism of solute transport and methods to study the same. The students are already exposed to laws of thermodynamics in the lower level however, they should be made aware of the electron transport chain in Prokaryotes and Mitochondria. ATP synthesis and anabolic mechanisms need to be explained to the students to understand the breakdown of mono, di- and oligosaccharides. The students will also be exposed to the fermentative pathways and anabolic reactions.

- Understand the architecture of the membrane and how solute is transported inside the cell.
- Describe and explain the electron transport chains in prokaryotes and mitochondria and understand the mechanism of ATP synthesis.
- Explain bioluminescence mechanism and its significance
- Discuss the experimental aspect of studying catabolism and anabolism and the various pathways for the breakdown of carbohydrates along with reactions in amphibolic pathways.
- Describe various other pathways which produce different end products.
- Describe anabolic reactions in carbohydrate synthesis.
- Apply the concepts of energetics and catabolism in biodegradation of various substrates.

Course Code	Title	Credits
RUSMIC503	MICROBIAL BIOCHEMISTRY PART I	2.5 Credits(65 lectures)
Unit I	BIOLOGICAL MEMBRANES & TRANSPORT	15 lectures
	1.1 Composition and architecture of membrane 1.1.1 Lipids	02
	1.1.2 Integral & peripheral proteins & interactions with	
	lipids	
	1.1.3 Permeability and outer membrane- a barrier	\mathbf{C}
	1.1.4 Aquaporins 1.1.5 Mechanosensitive channels	
	1.2 Methods of studying solute transport	02
	1.2.1. Using whole cells	
	1.2.2. Using Liposomes 1.2.3. Using Proteoliposome	
	1.2.3. Using Proteoliposonie	08
	1.3 Solute transport across membrane	
	1.3.1. Passive transport facilitated by membrane proteins.	
	1.3.2. Transporters grouped into Superfamilies'	
	1.3.3. Co transport across plasma membrane (Uniport, Antiport, Symport)	
	1.3.4. Active transport & electrochemical gradient	
	1.3.5. Ion gradient provides energy for secondary Active	
	transport e.g. Lactose transport	
	1.3.6. ATPases and transport	
	1.3.7. ABC transporters e.g. Histidine transport 1.3.8. Shock sensitive system - Role of binding proteins	
	e.g. Maltose uptake	
	1.3.9. Phosphotransferase system	
	1.3.10. Schematic representation of various	
	Membrane transport mechanisms in. E. coli	
	1.4 Other examples of solute transport-	
	1.4.1. Iron transport: A special problem	
	1.4.2. Bacterial protein export	03
NY.	1.4.3. Bacterial membrane fusion central to many	
	biological processes	
Unit II	BIOENERGETICS AND BIOLUMINESCENCE	15 lectures
	2.1. Biochemical mechanism of generating ATP-	01
	Substrate level, Oxidative, and Photo Phosphorylation	
	2.2. Electron transport chain	03
	2.2.1. Universal Electron acceptors that transfer Electro ETC.	

oteins, Quinones	
nur proteins, Cytochromes	
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IETC	
of Mitochondrial ETC	
ron carriers in bacteria 03	
ansport pathway in	
ases	
in <i>E. coli</i> - aerobic and	
in Azotobacter	
Proton motive force, Proto 04	
tio, Redox potential	
uring electron transfer from	
Mitochondrial ATP syntha	
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TP synthase	
itors of ATPase, Uncouple	
n of electrochemical	
02	
e Uzactate efflux	
tion, Significance, Function	
ent systems 02	
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YING METABOLISM & 15 lectures	Unit III
netabolism 03	
ch metabolism is studied.	
CARBOHYDRATES	Unit III

		1. Metabolic probes		
		2. Use of radioisotopes in biochemistry		
		i. Pulse labeling		
		ii. Assay & study of radiorespirometry -to differentiate EMP & ED		
		5. Use of biochemical mutants.		
		Sequential induction technique		
		3.2. Catabolism of Carbohydrates		5
		3.2.1. Breakdown of polysaccharides - glycogen, starch		
		cellulose.		
		3.2.2. Breakdown of oligosaccharides- lactose,		
		maltose, sucrose, cellobiose	10	
		3.2.3. Utilization of monosaccharides - fructose, Galact	\mathbf{O}	
		3.2.4. Major pathways-		
		i. Glycolysis (EMP)		
		ii. HMP Pathway & Significance of the pathway		
		iii. ED pathway,		
		iv. TCA cycle & Significance of the cycle	01	
		v. Anaplerotic reactions	UI	
		vi. Glyoxylate bypass,		
		vii.Incomplete TCA in anaerobic bacteria		
		viii.Amphibolic role of EMP and TCA cycle		
		ix.Energetics of Glycolysis, ED and TCA	04	
		Balance sheet and efficiency calculation	01	
	0	AMRUIAN		
	Unit IV	FERMENTATIVE PATHWAY& ANABOLISM OF CARBOHYDRATES	15 lectures	
8	M	1.Fermentative pathways (With structures and enzyme 4.1.1. Lactic acid fermentation - i. Homofermentors ii. Heterofermentors iii. Bifidobacterium pathway (Schematic)	04	
		4.1.2. Alcohol fermentation		
		i. by ED pathway in bacteria		



- 1. Stanier, R. Y.,M.Doudoroff and E. A. Adelberg. General Microbiology, 5th edition, The Macmillan press Ltd
- 2. Conn, E.E., P. K.Stumpf, G.Bruening and R. Y.Doi. 1987. Outlines of Biochemistry, 5th edition, 1987. John Wiley &Sons. New York.
- 3. Gottschalk, G., (1985), Bacterial Metabolism, 2nd edition, Springer Verlag
- 4. White, D., (1995), The Physiology and Biochemistry of Prokaryotes, 3^d edition, Oxford University Press
- Nelson, D. L. and M.M. Cox(2005), Lehninger, Principles of biochemistry.
 4th edition, W. H. Freeman and Company
- 6. Rose, A.H. (1976) Chemical Microbiology, 3^{re}dnButterworth-Heinemann
- 7. Zubay, G. L (1996), Biochemistry, 4th edition, Wm. C. Brown publishers
- 8. Mathews, C.K., K.E. van Holde, D.R. Appling, S,J, Anthony-Cahill (2012) Biochemistry, 4[∞]edn. Pearson
- 9. Wilson and Walker , 4^{edn}
- 10. Madigan, M.T. and J.M. Martinko 2006. Brock Biology of Microorganisms. Pearson Prentice Hall;
- 11. Zubay, G. L (1996), Principles of Biochemistry, Wm. C. Brown publishers
- 12. Cohen, G.N. (2011). Microbial Biochemistry. 2ndedn, Springer

Course Code: RUSMIC504 Course Title: BIOPROCESS TECHNOLOGY Academic year 2019-20

Learning Objectives

AMMAF

Bioprocess Technology course is designed to develop thelearner's ability to study the techniques used in the different phases of industrial microbiologysuch as strain improvement, basic fermentation equipment & its sterilization aspects. It gives an in-depth focus of the different types of fermenters used in industry for production of different products, and emphasizes its process parameters. It includes the principles and describes themain steps and processes in the industrial production of beverages and enzymes. The downstreamprocess and the environmental aspects of the final product are also included.

The last unit appraises the learner with instrumental techniques used in industry for analysis of products or intermediates during product development/ during fermentation/ during purification. It also introduces the learner to the legal aspects associated with fermentation industry in the form of Intellectual property rights.

- Describe the applications of microbes and its strain improvement in Industrial Microbiology.
- Apply kinetic formula to determine growth and productivity parameters of batch
 and continuous fermentations
- Describe the design of bioreactors for different applications and its process parameters
- Design media, growth conditions and techniques for producing and recovering different types of products of commercial value
- Design an industrial process by keeping in view the strict guidelines for its recovery &disposal
- Learner will be well -versed with the environmental aspects such as effluent treatment and carbon credits.
- Understand principle of working of important instruments used in biochemical, microbiological analysis.
- Get an overview of IPR and types of IP

Course Code	Title	Credits
RUSMIC504	BIOPROCESS TECHNOLOGY	2.5 Credits(65 lectures)
Unit I	UPSTREAM PROCESSING	15 lectures
	1.1: Strains and Strain Improvement of industrial microorganisms	10
	 i. Isolation of industrially important microorganisms ii. Improvement of industrial microorganisms a. Selection of induced mutants for primary metabolite b. Isolation of induced mutants for secondary metabolites. 1.2: Sterilization i.Introduction. Media sterilization (Concept of nabla factor) ii. Design and methods of batch sterilization iii. Design and methods of continuous sterilization 	5
Unit II	FERMENTER EQUIPMENT AND CONTROL	15 lectures
	 2.1.Design of fermenter i. Inoculum development ii.Basic functions of fermenter- Aseptic operation & containment, Body construction, Aeration and agitation iii.Achievement & maintenance of aseptic condition, Valves / Steam traps - function in general & examples. iv.Types of fermenters: Acetator, Cavitator, Tower fermenter, Cylindro conical, Air lift - outer loop / inner loop, Deep jet, Cyclone column, Packed tower (generator), Rotating disc, Bubble cap 	10
MAY	2.2: Control of variables Introduction, Types of sensors, Sensing & Control of- pH, temp, Dissolved oxygen, Flow measurement &control, Pressure, Inlet / Exit gas analysis, Foam sensing, oxygen	5
Unit III	DOWNSTREAM PROCESSING	15 lectures
	3.1. Downstream processing	10

	 Introduction, Precipitation, Filtration - theory, filter-aids, batch filters(Plate and frame filters), continuous filters(Rotary vacuum), Centrifugation: flocculating agent, range of centrifuges - Basket, tubular bowl. i. Cell disruption: Physico-chemical. ii. Liquid - Liquid extraction, Solvent recovery, iii. Chromatography -lon exchange &Adsorption Membrane processes - Ultrafiltration, reverse osmosis, liquid membranes. Drying, Crystallization, Whole broth processing. 3.2.1 Effluent treatment and regulations for fermentation industry 3.2.2. Modern methods of effluent treatment 3.2.3.Carbon Credits 	005	
Unit IV	BIOINSTRUMENTATION AND IPR	15 lectures	
	 4.1. Bioinstrumentation - Principles, working and applications of: i. Spectrophotometry (I. R) ii. Atomic absorption (AAS) & Atomic Emission 	8	
	 spectroscopy (Flame photometry) iii. Mass Spectroscopy- MALDI ToF, ESI 4.2. Intellectual Property Rights: Introduction to Intellectual Property Genesis of IPR - GATT, WTO, TRIPS, World Intellectual Property Rights Organization (WIPO) iii. Types of Intellectual Property - Patents, Copyright, Trademark, Trade secret, Plant varieties protection act, Designs, Geographical Indications 	7	

- Casida L. E., "Industrial Microbiology∥(2009) Reprint, New Age International (P)Ltd,Publishers, New Delhi
- 2. Stanbury P. F., Whitaker A. &Hall--S. J., (1997), "Principles of Fermentation Technology", 2nd Edition, Aditya Books Pvt. Ltd, New Delhi.
- 3. H. A. Modi, (2009). "Fermentation Technology" Vols 1 & 2, Pointer Publications, India
- 4. Okafor Nakuda (2007) 'Modern Industrial Microbiology and Biotechnology", Science Publications Enfield, NH, USA.

- 5. G Y Shitole and Ram Sable (2012) Environmental Degradation Issues And Challenges (Research publication)
- 6. Crueger W. and Crueger A. (2000) "Biotechnology -"A Textbook of Industrial Microbiology", 2nd Edition, Panima Publishing Corporation, New Delhi.

Practicals (Semester-V) RUSMICP501

FOR

[Practicals Based on 501, Credits -1.5 Lectures- 60]

- 1. UV survival curve determination of exposure time leading to 90% reduction
- 2. Isolation of mutants using UV mutagenesis
- 3. Replica plate technique for selection & characterization of mutants auxotroph & antibiotic resistant
- 4. Isolation and detection of plasmid DNA.
- 5. Preparation of competent cells and transformation
- 6. Demonstration of conjugation.

[Practicals Based on 502,Credits -1.5,Lectures-60]

- 1. Assignment on sample collection, transport and processing of any one pathological sample
- 2. Rapid Direct tests for identification of pathogens
 - a. Acid fast staining of *M. tuberculosis/ M.leprae*.
 - b. Metachromatic granule staining for C. diphtheriae
 - c. Catalase test
 - d. Bile solubility test
 - e. Slide coagulase test for S.aureus
 - f. Spot indole test
 - g. Oxidase test
 - h. Modern methods for identification of pathogens.
- 3. Identification of isolates obtained from following samples by morphological, cultural and biochemical properties
 - Nasal/ throat swabs(URT infection)
 - a. Sputum (LRT infection)
 - b. Skin swab/ pus (Skin infection)
 - c. Identification of $\ensuremath{\textit{Candida}}$ species using the germ tube test and growth on Chrom agar
 - d. Stool (GI tract infection)
 - e. Urine (UTI infection).
- 4. Demonstration of malarial parasite in blood film

- 5. Selection and testing of antibiotics using the Kirby-Bauer method
- 6. Determination of MIC of an antibiotic by E-test
- 7. Synergistic action of two drugs
- 8. Determination of MBC of an antibiotic.
- 9. Detection of βlactamase in S.aureus.
- 10. Role of plasmids in antibiotic resistance through curing of the plasmid

Practicals (Semester-V) RUSMICP502

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[Practicals Based on 503;Credits-1.5,Lectures- 60]

- 1. Isolation and detection of Mitochondria
- 2. Isolation and study of Bioluminescent organisms
- 3. Study of oxidative and fermentative metabolism
- 4. Carbohydrate fermentation tests
- 5. Mixed acid fermentations- Detection of organic acids by TLC
- 6. Study of Homo and Heterofermentation in Lactic acid bacteria
- 7. Detection of enzyme phosphatase
- 8. Quantitative assay of Phosphatase

[Practicals Based on 504, Credits -1.5, Lectures - 60]

- 1. Strip Plate Technique
- 2. Streak Plate Technique
- 3. Gradient plate technique for isolation of mutants.
- 4. Production and detection of vitamin B12 by bioautography.
- 5. Anaerobic digestion of Industrial effluent- Generation and detection of methane
- 6. Demonstration of IR spectroscopy and analysis of IR spectrum of one compound
- 7. Demonstration of GC-MS/ LC-MS.

AMMAR

Course Code: RUSMIC601 Course Title: GENE MANIPULATION, BIOINFORMATICS, CELL BIOLOGY &VIROLOGY Academic year 2019-20

Learning Objectives

This course introduces the learner to gene manipulation techniques which are an essential tool for modern day Genetic studies. This course also gives students theoretical and hands-oncompetence in major analytical techniques used inbioinformatics.

The section on Cell biology, although repeats some topics covered in FYBSc, is essentially to help the learner strengthen the basics of prokaryotic and eukaryotic cell structure. As the course on Biochemistry already deals with structure and function of cell membrane, the unit on Cell biology here does not repeat it.

Under the section of Virology, the course covers basic structure, life cycle of different types of viruses, genetics of lambda and cultivation of viruses. The course elaborates on different terminologies like cancer, prions, viriods and their mechanism.

- Understand fundamentals of gene manipulation
- Use bioinformatics tools for genetic analysis and structure building
- Correlate structure and function of important cell components of prokaryotic and eukaryotic cells
- Understand the basic structure, classification, enumeration, cultivation and life cycle of viruses
- Understand the terms like cancer, prions, viriods and their mechanisms
- Understand regulation of lambda phage

AMMARAMAR

Code
RUSMIC601
Unit I

CELL BIOLOGY 2.1 Structure and function of Prokaryotic cell a. Cell wall b. Capsule c. Flagella d. Endospore 2.2 Cytoskeleton and cell motility in eukaryotes	15 lectures 07 08
 a. Cell wall b. Capsule c. Flagella d. Endospore 	
 b. Capsule c. Flagella d. Endospore 	08
c. Flagellad. Endospore	08
d. Endospore	08
	08
2.2 Cytoskeleton and cell motility in eukaryotes	
 Cytosol, Ergastoplasm and cytoskeleton 	
b. Structure and function: Microtubules,	\bigcirc
Microfilaments, Intermediate filaments	
c. Microtubular organelles - Cilia, Flagella and	
BASIC VIROLOGY	15 lectures
3.1. Viral architecture-	04
1.1. Capsid, viral genome and envelope	
1.2. b. Structure of TMV, T4, Influenzavirus, HIV.	02
3.2. Viral classification	UZ
3.3 The viral replication cycle, attachment, penetration	04
	05
3.4. Cultivation of viruses- cell culture techniques,	00
embryonated egg, laboratory animals, Cell culture	
methods:Equipment required for animal cell	
culture, isolation of animal tissue	
ADVANCED VIROLOGY	15 lectures
4.1. Life cycle of T4 phage, TMV, Influenza Virus and	
HIV in detail	
4.0 Vieweliestien and enumeration of simo particles	
•	
,	
	 c. Microtubular organelles - Cilia, Flagella and centrioles d. Microfilament structures and role of associated proteins e. Molecular motors: Myosins, Kinesins, Dyenin BASIC VIROLOGY 3.1. Viral architecture- 1.1. Capsid, viral genome and envelope 1.2. b. Structure of TMV, T4, Influenzavirus, HIV. 3.2. Viral classification 3.3. The viral replication cycle- attachment, penetration, uncoating, types of viral genome and their replication, assembly, maturation and release. 3.4. Cultivation of viruses- cell culture techniques, embryonated egg, laboratory animals, Cell culture methods:Equipment required for animal cell culture, Isolation of animal tissue ADVANCED VIROLOGY 4.1. Life cycle of T4 phage, TMV, Influenza Virus and HIV in detail 4.2. A. Measurement of infectious units i. Plaque assay ii. Fluorescent focus assay

		-
v. Endpoint dilution assay.		
4.2.b. Measurement of virus particles and their		
components		
i. Electron microscopy		
ii. Atomic force microscopy		
iii. Haemagglutination		
iv. Measurement of viral enzyme activity.	, (^
4.3. Regulation of lytic and lysogenic pathway of		
lambda phage		
	\sim	
4.4. Role of viruses in cancer: Imp Definitions,	V	
characteristics of cancer cell, cancer multi step process,)	
Human DNA tumor viruses- EBV, Kaposis sarcoma virus,		
Hepatitis B and C virus, Papiloma Virus.		
4.5. Prions and viroids	02	

- 1. R. H. Tamarin, (2004), Principles of genetics, Tata McGraw Hill.
- 2. M.Madigan, J.Martinko, J.Parkar, (2009), Brock Biology of microorganisms∥, 12^e ed., Pearson Education International.
- 3. Fairbanks and Anderson, (1999), Genetics , Wadsworth Publishing Company.
- 4. Prescott, Harley and Klein, Microbiology 7th edition McGraw Hill international edition.
- 5. Edward Wagner and Martinez Hewlett, (2005) Basic Virology∥, 2nd edition, Blackwell Publishing
- 6. Teri Shors, (2009), Understanding viruses Jones and Bartlett publishers.
- 7. S.Ignacimuthu, (2005), Basic Bioinformatics∥, Narosa publishing house.
- 8. Robert Weaver, (2008), Molecular biology, , 3rd edn. McGraw Hill international edition.
- 9. Primrose and Twyman, (2001), Principles of gene manipulation and genomics∥, 6ed, Blackwell Publishing_
- 10. Arthur Lesk, (2009), Introduction to Bioinformatics , 3rd Edition, Oxford University Press
- 11. Snustad, Simmons, Principles of genetics , 3^{ee}dn. John Wiley & sons, Inc.
- 12. A textbook of biotechnology R.C.Dubey 4 *ed.S.Chand.
- 13. P.S Verma & V.K.Agarwal, ^CCell biology,genetics,molecular biology,evolution & ecology, 14th edn.
- 14. Lodish, Scott. Molecular cell biology, 7th edn, Macmillan higher education, international edn.
- 15. Ramsden Jerry,∥ Bioinformatics: An introduction, Springer international edition.
- 16. Flint, Enquist, Racanillo and Skalka, Principles of virology∥, (2009)3rdedn. ASM press
- 17. T. K. Attwood & D. J. Parry-Smith, (2003), Introduction to bioinformatics∥, Pearson education
- 18. Benjamin Lewin, (2014) 9th edition, [−]Genes IX∥, Jones and Bartlett publishers.
- 19. JD Watson, Baker (2004) 5thedn. Molecular biology of the gene∥, CSHL Press and Benjamin Cummings

Course Code: RUSMIC 602 Course Title: IMMUNOLOGY Academic year 2019-20

Learning objectives:

The course will help students to build on the basic information regarding Innate Immunity and Host Defence mechanisms that they have gained in S.Y. B.Sc. Immunology is an integral part of Medical Microbiology and this course is designed to help students understand the ability of our immune system to defend against invading pathogens in a logical fashion. This includes our innate ability to defend against microorganisms (innate immunity); should this first line of defense fail, how we can fight infections (acquired immunity). The course elaborates on the mechanisms of acquired defense after an introduction on the molecular nature of antigens and antibodies along with the role of different cells and their surface molecules in acquired immunity. After a basic introduction to cells of immune mechanisms the other units include details of mechanisms of acquired immunity- Humoral and Cell mediated. The curriculum also deals with how immune systems can fight infections (acquired immunity); if we react excessively, what price we pay (hypersensitivity); and very importantly, can we protect ourselves from diseased state (vaccination).

Learning Outcomes:

Students should be able to-

- Conceptualize how the innate and adaptive immune responses coordinate to fight
- invading pathogens
- Discuss the role of antigen in initiating the immune response
- Correlate the structure & functions of immunoglobulin
- Understand the importance of all the other entities involved i.e. T cells, B cells, NK cells, APCs, Cytokines, MHC, TcR, BcR, Co-receptors, Signalling pathways etc.
- Understand the effector responses- Humoral Immunity & Cell Mediated Immunity and differentiate between them
- Acquire an understanding of the role of immune system in disease: Unregulated response resulting in Hypersensitivity
- Understand the mechanism of Antigen-Antibody interaction & it's significance in diagnosis
- Apply the concept of immunity for protection from disease by development of vaccines

Course Code	Title	Credits
RUSMIC602	IMMUNOLOGY	2.5 Credits(65 lectures)
Unit I	ANTIGENS, ANTIBODIES AND ANTIGEN PRESENTATION	15 lectures
	1.1 : Antigens	05
	1.1.1 : Immunogenicity versus antigenicity	\mathbf{O}^{+}
	1.1.2: Factors that influence immunogenicity -)
	foreignness, molecular size, chemical composition, heterogenicity, ability to be processed and presented, contribution of the biological system to immunogenicity - genotype of the recipient, animal, immunogen dosage, route of administration and adjuvants	~
	1.1.3: Epitopes / antigen determinants (only concepts)	
	1.1.4: Haptens and antigenicity	
	1.1.5: Immunogenicity of some natural substances – native globular proteins, polysaccharides, lipids, nucleic acids Types of antigens - heterophile antigens, isophile antigens, sequestered antigens, super antigens, bacterial and viral antigens	
	1.2: Immunoglobulins	
	1.2.1 : Immunoglobulins - basic and fine structure 1.2.2: Immunoglobulin classes and biological activities	
	 1.2.3:Antigenic determinants on immunoglobulins - isotypes, allotypes, idiotypes 1.2.4 : Immunoglobulin Superfamily 	07
	1.2.5 : Monoclonal antibodies, Production	
	(Diagrammatically) & applications	
A,	1.3: T Cells, B cells and NK Cells	
<i>"C'</i>	1.4: Antigen presenting cells	
θ_{ll}	Antigen presentation- professional and nonprofessional cells and processing pathways,	01
	(Cytosolic and Endocytic pathway)	02
Unit II	ACTIVATION OF IMMUNE CELLS	15 lectures

	2.1 Cytokines	02
	Properties and functions Cytokines secreted by Th1 and Th2 cells	
	2.2 MHC complex and MHC molecules Structure of class I, and class II molecules; class III molecules Peptide - MHC interaction	03
	 2.3 T cells Receptors, structure (alpha-beta, gamma-delta TcR) TcR-CD3 complex structure & functions. Accessory molecules. Subsets of T cells (Th1, Th2, T reg) T cell activation, Costimulatory molecules, T cell differentiation (memory & effector cell) 	05
	2.4 B cells Receptors structure & organization	05
	 B cell activation and differentiation - i. Thymus dependent and independent antigens ii. B cell activating signals iii. Role of Th cells in Humoral response, formation of T - B conjugates, CD40 / CD40L interaction, Th cell cytokine signals. 	
Unit III	IMMUNE RESPONSES AND THEIR DETECTION	15 lectures
	 3.1.Humoral Response 3.1.1.Introduction of Humoral response, Primary and secondary responses 3.1.2.Germinalcentres and antigen induced B cell differentiation 3.1.3.Affinity maturation and somatic hyper mutation, Ig diversity, class switching 3.1.4.Generation of plasma cells and memory cells 	05
MAR	3.2. Cell mediated effector response 3.2.1. Generation and target destruction by Cytotoxic T cells.	03
	3.2.2. Killing mechanism of NK cells. 3.2.3. Antibody dependent cell cytotoxicity (ADCC)	06
		00
	3.3. Antigen-Antibody reactions	00

	agglutination inhibition, Radioimmunoassay (RIA), Enzyme immunoassays (EIA), Immunofluorescence, western blot technique 3.4.Immunodiagnostics Modern immunology based diagnostic tests	01
Unit IV	VACCINES, IMMUNOHEMATOLOGY AND HYPERSENSITIVITY	15 lectures
	 4.1: Vaccines 4.1.1 Active and passive immunization 4.1.2 Types of vaccines - Killed and attenuated vaccines, Whole organism vaccines, Purified macromolecules as vaccines, recombinant viral vector vaccines, DNA vaccines 4.1.3 Use of adjuvants in vaccine 4.1.4 New vaccine strategies, Ideal vaccine 4.2.1: Human blood group systems. ABO sporetors and 	05 05
	 4.2.1: Human blood group systems, ABO, secretors and non-secretors, Bombay Blood group. 4.2.2: Rhesus system and list of other blood group systems. 4.2.3: Haemolytic disease of new born, Coombs test. 4.3: Hypersensitivity Coombs and Gell's classification Type I to Type IV hypersensitivity, Mechanism and manifestation. 	05

- 1. Kuby Immunology, 6^a Edition, W H Freeman and Company
- 2. Pathak &Palan, Immunology: Essential & Fundamental, 1^e& 3^e Edition, Capital Publishing Company
- 3. Fahim Khan, Elements of Immunology, Pearson Education
- 4. Kuby Immunology, 7^h Edition, W H Freeman and Company

5. http://www.macmillanlearning.com/catalog/static/whf/kuby/

Course Code: RUSMIC603 Course Title: MICROBIAL BIOCHEMISTRY PART II Academic year 2019-20

Learning objectives:

There are a large number of macromolecules such as lipids, carbohydrates, proteins andnucleic acids which are catabolised by the living cells. Cells also bring about biosynthesis ofthese macromolecules. Various enzymes play a major role in these biochemical reactions. After an elaborate discussion on carbohydrate metabolism in the Semester V, the learner is made aware of the mechanisms of catabolism, anabolism as well as the regulation of lipid and nitrogenous compounds in this section.

Regulation of enzymatic reactions is a very critical part of metabolism. Studying these at the genetic level, would help students to get an insight on key mechanisms of economizing in the cells. Pathways for photosynthesis, with emphasis on prokaryotes are also dealt with here. Prokaryotic cells are alsoinvolved in metabolism of inorganic compounds. The last section elaborates on these mechanisms.

- Understand the reactions involved in metabolism of lipids and hydrocarbons.
- Describe and explain protein catabolism as well as anabolic processes in the cell.
- Explain nucleic acid metabolism and recycling of nucleotides.
- Discuss the mechanism of regulation with regards to allosteric proteins, gene expression as well as through other mechanisms like end product inhibition and covalent modification.
- Describe prokaryotic photosynthesis with respect to photosynthetic pigments, photochemical apparatus and light and dark reactions.
- Describe metabolism of inorganic compounds and Lithotrophy

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Course Code	Title	Credits
RUSMIC 603	MICROBIAL BIOCHEMISTRY PART II	2.5 Credits(65 lectures)
Unit I	PID METABOLISM & CATABOLISM OF HYDROCARBONS	15 lectures
	 1.1 General introduction to Lipids 1.1.1. Lipids and their functions 1.1.2. Action of lipases on triglycerides /tripalmitate 1.1.3. Phospholipids and their properties 1.1.4. Common phosphoglycerides in bacteria 	02
	 Catabolism of Lipids 1.2.1.Oxidation of saturated fatty acid- β oxidation pathway, Energetics of β oxidation of Palmitic acid 1.2.2. Oxidation of propionic acid. 1.2.3. Degradation of poly beta hydroxy butyrate 	05
	 Anabolism of Lipids Biosynthesis of straight chain even carbon saturated fatty acid (palmitic acid) Biosynthesis of phosphoglycerides in bacteria Biosynthesis of PHB 	06
	 Biosynthesis of PHB Catabolism of aliphatic hydrocarbons Oxidation of saturated aliphatic hydrocarbon (n-alkane) Omega oxidation pathway- Pathway in Corynebacterium and yeast Pathway in Pseudomonas 	02
Unit II	METABOLISM OF PROTEINS AND NUCLEIC ACIDS	15 lectures
MAR	 2.1 Protein catabolism 2.1.1. Enzymatic degradation of proteins 2.1.2. Metabolic fate of amino acids (schematic only) 2.1.3. Metabolism of single amino acids – Deamination reactions Decarboxylation Transamination 2.1.4. Fermentation of single amino acid -Glutamic acid by 	05
	<i>Clostridium</i> 2.1.5. Fermentation of pair of amino acids -Stickland	

	reaction	
	2. Amino acid synthesis	
	2.1. Schematic representation of amino acid families	
	ynthesis of amino acids of Aspartate family	04
	3. Nucleic acid Catabolism	
	1. Degradation of purine nucleotides up to	
	uric acid formation	03
	2. Recycling of purine and pyrimidine	
	nucleotides by salvage pathway	
	4. Anabolism of Nucleic Acids	\sim
	4.1. Metabolic origin of atoms in purine and pyrimidine ring)
	synthesis of pyrimidine nucleotides.	03
	synthesis of purine nucleotides.	
	rmation of deoxyribonucleotides.	
	.4.5. Synthesis of nucleotide diphosphates and	
	triphosphates.	
	le of nucleotides (high energy triphosphates)	
Unit III	METABOLIC REGULATION	15 lectures
	3.1: Overview and major modes of regulation	01
	Examples of cellular control mechanism acting at	
	various levels of metabolism (tabulation only)	
	2. Allosteric proteins	03
	3.2.1. Definition 3.2.2. Allosteric enzymes - Role of allosteric enzymes	
	using ATCase as example (no kinetic study)	
	3.2.3.Regulatory allosteric proteins	
	i. Interaction of proteins with DNA	
	ii. Structure of DNA Binding proteins	
	iii Everales Les represent Tra represent CAD protein	
	iii. Examples - Lac repressor, Trp repressor, CAP protein	
	iv.Definition and examples of alarmones	
	iv.Definition and examples of alarmones	
.0	iv Definition and examples of alarmones 3.3 Regulation of gene expression (Transcription)	
JP.	 iv.Definition and examples of alarmones 3.3 Regulation of gene expression (Transcription) 3.3.1. Introduction to operon model 	06
NA	iv Definition and examples of alarmones 3.3 Regulation of gene expression (Transcription)	06
MNA	 iv.Definition and examples of alarmones 3.3 Regulation of gene expression (Transcription) 3.3.1. Introduction to operon model 3.3.2. Common patterns of regulation of transcription -General concept of positive and negative regulation of operons i. <i>Lac</i> operon - Mechanism of regulation - Induction 	06
AMAR	 iv.Definition and examples of alarmones 3.3 Regulation of gene expression (Transcription) 3.3.1. Introduction to operon model 3.3.2. Common patterns of regulation of transcription -General concept of positive and negative regulation of operons i. <i>Lac</i> operon - Mechanism of regulation - Induction - Catabolite repression 	06
AWNE	 iv.Definition and examples of alarmones 3.3 Regulation of gene expression (Transcription) 3.3.1. Introduction to operon model 3.3.2. Common patterns of regulation of transcription -General concept of positive and negative regulation of operons i. <i>Lac</i> operon - Mechanism of regulation - Induction - Catabolite repression ii. <i>Trp</i> operon - End Product Repression 	06
AMAR	 iv.Definition and examples of alarmones 3.3 Regulation of gene expression (Transcription) 3.3.1. Introduction to operon model 3.3.2. Common patterns of regulation of transcription -General concept of positive and negative regulation of operons i. <i>Lac</i> operon - Mechanism of regulation - Induction - Catabolite repression 	06

	ii. Riboswitches	
	3.4 Regulation of enzyme activity (Post translational regulation) 3.4.1. End-Product Inhibition and Mechanism of End	04
	 Product Inhibition in branched pathways with examples a. Isofunctional enzymes b. Concerted feedback inhibition c. Sequential feedback inhibition d. Cumulative Feedback inhibition e. Combined activation and inhibition 3.4.2 Covalent modifications of enzymes i. General examples without structure ii. Monocyclic cascade &inter-convertible enzyme definition 	
	 ii. Glutamine synthetase system of E.coli 3.4.3. Regulation by proteolytic cleavage 3.5 Regulation of EMP and TCA (Schematic and Role of Pyruvate dehydrogenase Complex) 	01
Unit IV	PROKARYOTIC PHOTOSYNTHESIS & INORGANIC METABOLISM	15 lectures
	1. Prokaryotic photosynthesis	09
	 4.1.1. Early studies on photosynthesis Light and dark reactions Bacterial photosynthesis Hill reaction 2. Phototrophic prokaryotes -Oxygenic, Anoxygenic phototrophs examples only 3. Photosynthetic pigments 4. Location of photochemical apparatus 5. Photophosphorylation 6. Light reactions in i. Purple photosynthetic bacteria ii. Green sulphur bacteria 	
8 AMM	 iii. Cyanobacteria (with details) 7. Dark reaction i. Calvin Benson cycle ii. Reductive TCA 2. Inorganic Metabolism 4.2.1. Assimilatory pathways- i. Assimilation of nitrate, ii. Ammonia fixation - Glutamate dehydrogenase, Glutamine synthetase, GS-GOGAT, Carbamoyl 	03

iii. Biological nitrogen fixation (Mechanism for N₂fixation and protection of nitrogenase) iv Assimilation of sulphate
2. Dissimilatory pathways
i. Nitrate as an electron acceptor

(Denitrification in *Paracoccusdenitrificans*)
ii. Sulphate as an electron acceptor
4.2.3: Lithotrophy- Enlist organisms and products formed during oxidation of Hydrogen, carbon monoxide, Ammonia, Nitrite, Sulphur, Iron.
4.2.3: Lithotrophy- Enlist organisms and products formed during oxidation of Hydrogen, carbon monoxide, Ammonia, Nitrite, Sulphur, Iron.
4.2.3: Lithotrophy- Enlist organisms and products formed during oxidation of Hydrogen, carbon monoxide, Ammonia, Nitrite, Sulphur, Iron.

- 1. Stanier, R. Y., M. Doudoroff and E. A. Adelberg. General Microbiology, 5th edition, The Macmillan press Ltd
- Conn, E.E., P. K. Stumpf, G. Bruening and R. Y. Doi. 1987. Outlines of Biochemistry, 5th edition, 1987. John Wiley & Sons. New York.
- 3. Gottschalk, G., (1985), Bacterial Metabolism, 2nd edition, Springer Verlag
- 4. White, D., (1995), The Physiology and Biochemistry of Prokaryotes, 3^d edition, Oxford University Press
- Nelson, D. L. and M.M. Cox (2005), Lehninger, Principles of biochemistry.
 4th edition, W. H. Freeman and Company.
- 6. Salle, A.J. Fundamental Principles of Bacteriology, 7^eedn McGraw Hill Book Co.
- 7. Cohen, G.N. (2011). Microbial Biochemistry. 2^medn, Springer
- 8. Madigan, M.T. and J.M. Martinko 2006. Brock Biology of Microorganisms. Pearson Prentice Hall;
- 9. Biochemistry 3rd edition, Mathew, Van Holde and Ahern, Pearson Education
- 10. Zubay, G. L (1996), Biochemistry, 4th edition, Wm. C. Brown publishers
- 11. Principles of Biochemistry, Lehninger, 5^eednW. H. Freeman and Company

Course Code: RUSMIC604 Course Title: INDUSTRIAL MICROBIOLOGY Academic year 2019-20

Learning Objectives

The learner was introduced to fermentation technology in Semester V. This semester the learner is introduced to industrial fermentations for brewing, pharmaceutical and food industry. This section of the curriculum also includes traditional fermentation processes of alcoholic beverages and modern fermentations that acquaint the learner to exploit microbial. technology to make greener fuels. The learner is provided with thedetails of productions of important products like antibiotics, vitamins, organic acid, food products and supplementsandenzymes.

Bioassays as analysis techniques used by quality control or R & D labs of industries for various products are also dealt with here. The learner is expected to learn the need of Quality management as theproducts need to fulfil these requirements. Thus, this paper readies the learner to understand and apply the knowledge of fermentation technology and related products. This course aims to enable graduates to enter industry with an appropriate level of understandingof the need for both the science and business aspects to be achievable to make a viable product and enhance their entrepreneurial skills.

- Understand the actual process involved in fermentations of important beverages, pharmaceutical and food products.
- Learn the applications of enzymes in various fields.
- · Understand the principle of bioassays as an analytical technique
- Learn the salient features of quality management.

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Course Code	Title	Credits					
RUSMIC 604	INDUSTRIAL MICROBIOLOGY	2.5 Credits(65 lectures)					
Unit I	INDUSTRIAL FERMENTATIONS:	15 lectures					
	1.1. Types of alcoholic beverage.	V					
	1.2. Beer - Ale and Lager	3					
	1.3. Wine -Red and white & Champagne						
	1.4. Vinegar (acetator& Generator)	2					
	1.5. Bioethanol production-	3					
	-From feedstock to fermentable sugars	Ū					
	- Zymomonas mobilis as an alternate ethanol producer						
	1.6. Acetone Butanol Fermentation	2					
Unit II	II INDUSTRIAL FERMENTATIONS:II						
	2.1 Production of secondary metabolites-Antibiotics- Penicillin& Semisynthetic Penicillins	04					
	2.2 Production of primary metabolites-						
	i. Vitamin B ₁₂ from <i>Propionibacterium</i> & <i>Pseudomonas</i>	03					
	ii. Amino acids- Methods for manufacture, Glutamic	01					
	Acid (direct)	04					
	iii. Organic acids- Citric acid	01 05					
	iv. Enzymes- Uses of enzymes in industry, Production of Fungal amylase by solid substrate fermentation,	05					
	Stabilization of enzymes- Immobilization techniques						
	v. Biotransformation of steroids	01					
Unit III	INDUSTRIAL FERMENTATIONS: II	15 lectures					
	Mushroom cultivation	03					
	 SCP- Substrates used, Organisms and safety 	03					
	i.Fermented foods- Bread, fermented cassava, tea and	03					
\mathcal{O}	coffee Mold modified foods, Types (list only), Braduction of						
	. Mold modified foods- Types (list only), Production of Soya sauce	02					
	Lactic acid starter cultures, prebiotics and probiotics						
		04					

Unit IV	BIOASSAYS & QUALITY ASSURANCE	15 lectures	
	 4.1 Bioassays Comparison of Chemical and Biological assays Microbiological assays- Test organisms,types of assay methods and factors affecting. Modern methods for assay of fermentation products 	05	5
	 4.2 QA, QC, GMP: Definitions- Manufacture, Quality, Quality Control, In- Process Control, Quality Assurance, Good Manufacturing Practices. Chemicals & Pharmaceutical production: The five variables, In process Items, Finished Products, Labels and Labelling, Packaging materials, Documentation, Regulations. Control of Microbial contamination during manufacture: Premises and contamination control 	07	
	 Manufacture of sterile products, Clean and Aseptic Area, Important publications related to QA 4.3 Sterilization Control and Sterility Assurance: Bio-burden determinations Environmental monitoring Sterilization Monitors - Physical, Chemical and Biological indicators Sterility Testing 	03	

- 1. Casida L. E., "Industrial Microbiology∥ 2009 Reprint, New Age International (P) Ltd, Publishers, New Delhi
- 2. Crueger W. and Crueger A. 2000 "Biotechnology -"A Textbook of Industrial Microbiology", 2nd Edition, Panima Publishing Corporation, New Delhi.
- 3. H. A. Modi, 2009. 'Fermentation Technology' Vol: 1 & 2, Pointer Publications, India
- 4. Prescott and Dunn's 'Industrial Microbiology" (1982) 4th Edition, McMillan Publishers
- 5. Hugo & Rusell's, Pharmaceutical Microbiology Blackwell Science, Seventh Edition
- 6. Peppler, H. J. and Perlman, D. (1979), "Microbial Technology". Vol 1 & 2, Academic Press.
- 7. Michael J. Waites, 2001 Industrial Microbiology: An Introduction∥, Blackwell Science Publications
- 8. Naduka Okafor, Modern Industrial Microbiology , Science Publications, 2007

Practicals (Semester-VI) RUSMICP602

[Practicals Based on 603;Credits -1.5,Lectures- 60]

- 1. Qualitative detection of Lipase
- 2. Estimation of proteins by Lowry's method
- 3. Qualitative detection of Protease
- 4. Assay of enzyme Protease
- 5. Study of breakdown of amino acids Lysine decarboxylase and Deaminase activity

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- 6. Estimation of uric acid
- 7. To study catabolite repression
- 8. Study of Hill reaction
- 9. Study of photosynthesis in cyanobacteria
- 10. Study of Lithotrophs Nitrosification and Nitrification

[Practicals Based on 604;Credits:1.5, Lectures:60]

- 1. Alcohol tolerance for yeast.
- 2. Sugar tolerance for yeast.
- 3. Inoculum Development for alcohol fermentation
- 4. Alcohol fermentation .: Efficiency of fermentation
- 5. Chemical estimation -Sugar by Cole's Ferricyanide method
- 6. Chemical estimation -Alcohol Estimation Dichromate method
- 7. Demonstration of HPLC for alcohol estimation
- 8. Production of fungal amylase using solid substrate fermentation
- 9. Immobilization of yeast invertase
- 10. Mushroom cultivation
- 11. Production of Spirulina SCP
- 12. Isolation of Lactic acid bacteria from probiotics
- 13. Bioassay of an antibiotic (Ampicillin / Penicillin)
- 14. Bioassay of Cyanocobalamin.
- 15. Chemical estimation of Penicillin
- 16. Sterility testing of water for injection or DPT vaccine.

Modality of Assessment:

Theory Examination Pattern:

A. Internal Assessment - 40% 40 marks.

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	Evaluation type	Marks
	One Assignment/Case study/Project	10
	One class Test (multiple choice questions / objective)	20
	Active participation in routine class instructional deliveries(case studies/ seminars//presentation)	05
	Overall conduct as a responsible student, manners, skill in articulation, leadership qualities demonstrated through organizing co-curricular activities, etc.	05

B) External examination - 60 %

Semester End Theory Assessment - 60%

60 marks

1. Duration - These examinations shall be of **two hours** duration.

2. Theory question paper pattern: -There shall be **four** questions each of **15** marks, one on each unit. All questions shall be compulsory with internal choice within the questions.

Paper Pattern:

Questions	Options	Marks	Questions on
Q.1)A)	Any 2 out of 4	12	Unit I
Q.1)B)	Any 3 out of 5	03	
Q.2)A)	Any 2 out of 4	12	Unit II
Q.2)B)	Any 3 out of 5	03	
Q.3)A)	Any 2 out of 4	12	Unit III
Q.3)B)	Any 3 out of 5	03	
Q.4)A)	Any 2 out of 4	12	Unit IV
Q.4)B)	Any 3 out of 5	03	

Practical Examination Pattern:

(A) Internal Examination: -

	Paper I	Paper II	Paper III	Paper IV	
Journal	05	05	05	05	
Tests	10	10	10	10	
Participation	05	05	05	05	
Total	20	20	20	20	

(B) External (Semester end practical examination):- 30 Marks Per Section

Sr. No.	Particulars	Marks	Total	
1	Lab work	50+50=	100	
2	Viva	10+10=	20	

PRACTICAL BOOK/JOURNAL

Semester V and VI:

The students are required to present a duly certified journal for appearing at the practical examination, failing which they will not be allowed to appear for the examination.

In case of loss of Journal and/ or Report, a Lost Certificate should be obtained from Head/ Co-ordinator / In charge of the department; failing which the student will not be allowed to appear for the practical examination.

Overall Examination and Marks DistributionPattern

Semester V

Course	501					502			503			504	Grand Total
	In	Ex	Total	In	Ex	Total	In	Ex	Total	In	Ex	Total	
Theory	40	60	100	40	60	100	40	60	100	40	60	100	400
Practical	20	30	50	20	30	50	20	30	50	20	30	50	200

Semester VI

7	Course		601 602 603								604	Grand Total		
		In	Ex	Total	In	Ex	Total	In	Ex	Total	In	Ex	Total	
	Theory	40	60	100	40	60	100	40	60	100	40	60	100	400
	Practical	20	30	50	20	30	50	20	30	50	20	30	50	200