# **VIVEKANANDHA**

# **COLLEGE OF ARTS AND SCIENCES FOR WOMEN**

[AUTONOMOUS]

(Affiliated to Periyar University, Approved by AICTE & Accredited by NAAC)

Recognised under section 2(f) and 12(B) of UGC Act, 1956

ELAYAMPALAYAM, TIRUCHENGODE (Tk.), NAMAKKAL (Dt.).



DEPARTMENT OF MICROBIOLOGY
B.Sc., MICROBIOLOGY
SYLLABUS & REGULATIONS

[FOR CANDIDATES ADMITTED FROM 2019-2020 ONWARDS UNDER AUTONOMOUS & CBCS PATTERN]

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VIVEKANANDHA EDUCATIONAL INSTITUTIONS Angammal Educational Trust Elayampalayam, Tiruchengode (Tk.), Namakkal (Dt.)

## **B.Sc.**, Microbiology

## 1. SCOPE OF THE COURSE

The course of Microbiology is intended to prepare the students not only to be knowledgeable in the science of Microbiology, but also to be useful in the upliftment of the social and economic well being. Courses offered cover all areas of basic and applied microbiology and these prepare students for a Bachelor of Science degree in Microbiology.

The degree is a three-year full time programme. The programme is not only a specialist programme, but it is also designed to be relevant to the social and economic needs of the nation. In reflection to the specialized nature of the programme, emphasis is given to practical and acquisition of practical skills.

The Programme has been involved in teaching basic and applied microbiology as well as making findings on local problems of microbiology interest. The vision of the programme is therefore, to produce graduates who are not only knowledgeable in the science of microbiology, but who can make significant contributions to the development the human society.

The programme is aimed at training undergraduate graduate students who would have adequate background knowledge and practical skills for application in postgraduate research, teaching, industrial production, medicine, environmental management and biotechnology.

## 2. SALIENT FEATURES

- ❖ Course is specially designed for a higher level career placement.
- Special guest lecture from industries will be arranged.
- \* Enables students to gain a job oriented degree.
- Special industry orientations and training are parts of the degree course.

## 3. OBJECTIVES OF THE COURSE

## The specific objectives of the programme are:

- ❖ To equip the undergraduate students with a sound knowledge of the fundamental principles involved in the study of microbiology.
- ❖ To produce graduates that would make impact in the diverse fields of human endeavor considering the ubiquitous nature of microorganism and the wide − ranging applications of the knowledge of microbiology.
- ❖ To provide focus for a career in various fields of applied science including medicine, pharmacy, bio-mining, biotechnology, industrial production, environmental management, agriculture and bioinformatics.

#### 4. ELIGIBILITY FOR ADMISSION

Candidates seeking admission to the first year degree course for **B.Sc.**, **Microbiology** shall be required to have passed

- a) Higher secondary examination with biology as major subjects conducted by the Government of Tamil Nadu (or)
- b) These regulations shall take effect from the academic year 2017-2018 i.e. for the students who are to be admitted to the first year of the course during the academic year 2017-2018 and thereafter
- c) Any examination with biology as major subjects of any other University or Board accepted as equivalent there to by Periyar University.
- d) Academic and vocational stream candidates are eligible.

## 5. DURATION OF THE COURSE

- The course shall extend over a period of three academic years consisting of six semesters. Each academic year will be divided into two semesters. The first semester will consist of the period from July to November and the second semester from December to March.
- The subjects of the study shall be in accordance with the syllabus prescribed from time to time by the Board of Studies of Vivekanandha College of Arts and Sciences for Women (Autonomous) with the approval of Periyar University.
- Each subject will have six hours of lecture per week apart from practical at the end of even semester.

#### 6. CONTINUOUS INTERNAL ASSESSMENT

The performance of the students will be assessed continuously and the Internal Assessment Marks will be as under:

## **Theory**

Average of two tests
 Assignment
 Marks
 Attendance
 Marks
 Marks

Total
25 Marks

## **Practical**

Practical best average of two tests - 30 Marks
 Attendance - 5 Marks
 Observation note - 5 Marks

Total 40 Marks

## **Break-up Details for Attendance**

Below 75% - No Marks

76 to 80% - 1 Mark

81 to 85% - 2 Marks

86 to 90% - 3 Marks

91 to 95% - 4 Marks

96 to 100% - 5 Marks

#### **PASSING MINIMUM**

## **INTERNAL**

There shall be no passing minimum for internal

#### **EXTERNAL**

In the end semester examinations, the passing minimum shall be 40 % out of 75 Marks (30 Marks)

## 7. ELIGIBILITY FOR EXAMINATION

A candidate will be permitted to appear for the end semester examination only on earning 75 % of attendance and only when his/her conduct has been satisfactory. It shall be open to grant exemption to a candidate for valid reasons subject to conditions prescribed.

## 8. CLASSIFICATION OF SUCCESSFUL CANDIDATES

Successful candidates passing the examination of language, core, allied, elective, skill based elective and non major elective courses and securing marks

- a) 75% and above shall be declared to have passed the examination in first class with Distinction provided they pass all the examinations prescribed for the course at first appearance itself.
- b) 60% and above but below 75% shall be declared to have passed the examinations in first class without distinction.
- c) 50% and above but below 60% shall be declared to have passed the examinations in second class.

- d) All the remaining successful candidates shall be declared to have passed the examinations in third class.
- e) Candidates who pass all the examinations prescribed for the course at the first appearance itself and within a period of three consecutive academic years from the year of admission only will be eligible for University rank.

#### 9. ELIGIBILITY FOR AWARD OF THE DEGREE

A candidate shall be eligible for the award of the degree only if she has undergone the above degree for a period of not less than three academic years comprising of six semesters and passed the examinations prescribed and fulfilled such conditions has have been prescribed therefore.

## 10. PATTERN OF QUESTION PAPER

**PART- A** (Objective) Answer all Questions  $20 \times 1 = 20 \text{ Marks}$ 

**PART- B** (500 words) Answer all 5 Questions (either or type)  $5 \times 5 = 25$  Marks

**PART - C** (1000 words) Answer any 3 Questions (three out of five)  $3 \times 10 = 30$  Marks

## 11. PROCEDURE IN THE EVENT OF FAILURE

If a candidate fails in a particular subject, she may reappear for the university examination in the concerned subject in subsequent semesters and shall pass the examination.

## 12. COMMENCEMENT OF THESE REGULATIONS

These regulations shall take effect from the academic year 2017 - 2018 i.e. for the students who are to be admitted to the first year of the course during the academic year 2017 - 2018 and thereafter.

#### 13. TRANSITORY PROVISION

Candidates who were admitted to the UG course of Microbiology before 2017 - 2018 shall be permitted to appear for the examinations under those regulations for a period of three years *i.e.*, up to and inclusive of the examination of April/May 2020. Thereafter, they will be permitted to appear for the examination only under the regulations then in force.

## Vivekanandha College

## VISION

To evolve into a centre of excellence in higher education through creative and innovative practices to secure social equity for women.

## **MISSION**

- 1. To provide sufficient learning infrastructure to the students to pursue their studies
- 2. To provide good opportunity for higher education and conducive environment to the students to acquire education
- 3. To provide high quality academic programme, training activities and research facilities
- 4. To facilitate industry-institute interface

## **VISION**

Aspires to be a microbiologist committed to progress the quality of human lives by exploring environment, fighting with disease and to utilize microbes for healthy food.

## **MISSION**

To educate the students to acquire the academic excellence with national and international recognition

To train the students to recognize, investigate and to resolve the myriad of microbiological problems affecting health and the environment through the programme designs

To contribute to the cutting edge in Microbiology by pursuing high quality research and other scholarly activities

To motivate the students to become a women entrepreneur by applying their knowledge in the field of microbiology

To establish as an expert resource within the geographical areas regarding all issues related to medical and environmental microbiology

## **B.Sc., MICROBIOLOGY**

## **PROGRAMME OUTCOME:**

The programme aims to communicate the scientific knowledge relating to microbiology and their role in the ecosystem and health issues. It is designed to teach and practice the fundamentals of microbiology, by experts in microbiology for the development of microbiology across the society.

## PROGRAMME SPECIFIC OUTCOME:

- 1. To describe about the basics of microbiology, genetics, metabolism and ecology.
- 2. To make the students understand the integration of microbes and their role in causing disease with the immune status of immune system in diagnosis and treatment.
- 3. To train them in the application of microbiology with the components of laboratory skills.
- 4. To explain the ubiquitous nature of microbes in terms of their wide range of ecological habitats.
- 5. To comprehend the effectiveness of microbes in biotechnology, fermentation technology, medicine and other industries for human welfare.

# $SCHEME\ OF\ CURRICULUM-B.Sc.,\ IN\ MICROBIOLOGY$ (For the candidates admitted during the academic year 2018 – 2019 onwards)

Sem	Subject code	Part	Course	Subjects	Hrs/ Week	Credits	Int. Marks	Ext. Marks	Tot. Marks
	18U1LT01			Tamil – I					
	18U1LH01	I	Language – I	Hindi – I	6	3	25	75	100
	18U1LM01			Malayalam – I					
	17U1LE01B	II	English – I		6	3	25	75	100
I	18U1MBC01	III	Core – I	Principles of Microbiology	5	5	25	75	100
	18U1MBCP01		Core	Major Practical – I	4	3	40	60	100
	181U1BCA01	777	A11' 1 T	Biochemistry	4	4	25	75	100
	18U1BCAP01	III	Allied – I	Allied Practical – I	3	2	40	60	100
	18U1VE01	IV		Value education – (Yoga)	2	2	25	75	100
				Total	30	22	205	495	700
	18U2LT02			Tamil – II					
	18U2LH02	I	Language – II	Hindi – II	6	3	25	75	100
	18U2LM02			Malayalam – II	1				
	18U2LE02B	II	English – II		6	3	25	75	100
II	18U2MBC02	III	Core – II	Microbial Physiology and Metabolism	4	4	25	75	100
	18U2MBCP02	III		Major Practical – II	3	2	40	60	100
	18U2MBA01	III	Allied – II	Bioinstrumentation Techniques	4	4	25	75	100
	18U2MBAP01	III		Allied Practical – II	3	2	40	60	100
	18U2ES01	IV		Environmental studies	4	4	25	75	100
				Total	30	22	205	495	700
	18U3LT03			Tamil – III	6			75	
	18U3LH03	I	Language – III	Hindi – III		3	25		100
	18U3LM03			Malayalam – III	1				
	17U3LE03B	II	English – III		6	3	25	75	100
	18U3MBC03	III	Core – III	Molecular Biology and Microbial Genetics	4	4	25	75	100
	18U3MBCP03			Major Practical – III	3	2	40	60	100
	18U3MBA02	III	Allied – III	Bioinformatics	4	4	25	75	100
III	18U3MBAP02	111	7 tined in	Allied Practical – III	3	2	40	60	100
	18U5CCN06			Interview Skills					
	18U5CCN01			Entrepreneurial					
			\.n.ma	Development					
	18U5TAN01	IV	NMEC – I	Pechukalai	2	2	25	75	100
	18U3CHN01			Industrial Chemistry	1				
	18U6CHE02 18U3ZON01			Medicinal Chemistry Sericulture	_				
	18U3MAAS01	IV	SBEC – I	Biostatistics	2	2	25	75	100
	TOUSWIAASUT	1 4	SBEC -1	Total	30	22	230	<b>570</b>	800
	18U4LT04			Tamil – IV				270	- 000
	18U4LH04	I	Language – IV	Hindi – IV	6	3	25	75	100
	100-1110-	1	Lunguage 1V		-	3		, 5	100
***	18H4LM04			I Malayalam – IV					
IV	18U4LM04 18U4LE04	II	English – IV	Malayalam – IV	6	3	25	75	100
IV	18U4LM04 18U4LE04 18U4MBC04	II	English – IV  Core – IV	Malayalam – IV  Immunology and	6	3 4	25 25	75 75	100

	18U4MBCP04			Major Practical – IV	3	2	40	60	100
	18U4BTA01	III	Allied – IV	Biotechnology	4	4	25	75	100
	18U4BTAP01	ш		Allied Practical – IV	3	2	40	60	100
		IV	NMEC – II	Elected by Students	2	2	25	75	100
	18U4MBS02	IV	SBEC – II	Plant Diseases and Management	2	2	25	75	100
				Total	30	22	230	570	800
	18U5MBC05	III	Core – V	Medical Bacteriology and Mycology	6	6	25	75	100
	18U5MBC06	III	Core – VI	Industrial and Pharmaceutical Microbiology	5	5	25	75	100
V	18U5MBC07	III	Core – VII	Genetic Engineering	5	5	25	75	100
	18U5MBE01/02	III	Elective – I	Elected By Students	4	4	25	75	100
	18U5MBS03	IV	SBEC – III	Computer Applications in Biology	2	2	25	75	100
	18U5MBMP01			Mini Project	2	1	-	ı	ı
	18U5MBCP05	III		Practical – V		3	40	60	100
				Total	30	26	165	435	600
	18U6MBC08	III	Core – VIII	Medical Virology and Parasitology	6	6	25	75	100
	18U6MBC09	III	Core – IX	Soil and Environmental Microbiology	5	5	25	75	100
VI	18U6MBC10	III	Core – X	Food and Dairy Microbiology	5	5	25	75	100
	18U6MBE03/04	III	Elective – II	Elected by Students	4	4	25	75	100
	18U6MBS04	IV	SBEC – IV	Advances in Microbiology	2	2	25	75	100
	18U6MBCP06	III		Practical – VI	6	3	40	60	100
	18U6MBEX01	-	-	Extension activity	2	1	-		
				Total	30	26	165	435	600
		180	140	1200	3000	4200			

## **MAJOR ELECTIVE COURSES:**

## Semester – V

- 1. Hematology and Blood Banking (18U5MBE01)
- 2. Entrepreneurship in Microbiology (18U5MBE02)

## Semester – VI

- 1. Microbial Diagnosis in Health Clinics (18U6MBE03)
- 2. Quality Control in Food Microbiology (18U6MBE04)

## NON MAJOR ELECTIVE COURSES:

- 1. Public Health and Hygiene (18U3MBN01)
- 2. Diseases Epidemics and Control (18U4MBN02)



**CORE - I** 

Total Number of Hours: 60 5 Hours/ Week

#### PRINCIPLES OF MICROBIOLOGY

## **Course Objectives:**

- To study the history and scope of Microbiology
- To gain knowledge about techniques in Microbiology
- To understand the cultivation techniques of microbes
- To study the classification of bacteria
- To gain knowledge on diverse group of bacteria

#### **Course Outcome:**

CO1	The students could understand the origin of Microbiology field and its discoveries in
	reference to the contributions of great scientists
CO2	The use of microscopy and the methods to visualize the microorganisms were could
	be learnt
CO3	The art of cultivating the microorganisms, storing methods and removal of pathogenic
	organisms were taught
CO4	The students could learn the diverse groups of microorganisms
CO5	The microorganisms that grow at some extreme conditions were to be introduced

UNIT – I No. of Hours: 12

**History and Development of Microbiology:** Spontaneous generation verses biogenesis. Contributions of Anton van Leeuwenhoek, Louis Pasteur, Robert Koch, Joseph Lister and Alexander Fleming – Germ theory of disease and golden era of microbiology. Contributions of Martinus W. Beijerinck, Sergei N. Winogradsky and Selman A. Waksman. Paul Ehrlich, Elie Metchnikoff and Edward Jenner. Scope of microbiology.

UNIT – II No. of Hours: 12

**Microscopy:** Bright field, Dark Field, Phase contrast and Fluorescence microscope. **Staining Methods:** Staining and its types – Simple staining, Differential staining – Gram's, Acid fast and Special staining methods – Endospore and Capsule staining. Hanging drop technique.

UNIT – III No. of Hours: 12

**Cultivation of Microbes:** Culture media and its types. Cultivation of anaerobes – Pyrogallol and Gas Pak method. Pure culture isolation techniques. **Sterilization:** Physical and Chemical methods of sterilization. Preservation of cultures. Antibiotics classification based on mode of action – Tests for sensitivity to antimicrobial agents.

UNIT – IV No. of Hours: 12

**Microbial Diversity:** Evolution, Phylogeny, Microbial Taxonomy and Classification – Haeckel, Whittaker and Carl Woese system, Numerical Taxonomy and Molecular based classification. Bacterial diversity – General characteristics of bacteria and classification – Bergeys' Manual of Systematic Bacteriology (up to order level) and Actinobacteria.

UNIT – V No. of Hours: 12

**General characteristics:** of Chlamydia, Rickettsia and Mycoplasma. Microbial diversity in different ecosystems - psychrophiles, mesophiles, thermophiles, acidophiles, alkalophiles, barophiles, capnophilic, saccharophilic and other extremophiles (Halophiles, Methanogens). Economic importance of bacteria.

## **Text Books**

- **1.** Pelczar MJ, Chan ECS and Kreig NR (2008). **Microbiology**. 5<sup>th</sup> Edition, Tata McGraw Hill Education Pvt. Ltd., New Delhi.
- **2.** Dubey RC and Maheswari DK (2013). **A Textbook of Microbiology.** 3<sup>rd</sup> Edition. S Chand and Company Limited, New Delhi.
- **3.** Sullia S.B and Santhanam S (2017). **General Microbiology.** 2<sup>nd</sup> Edition, Oxford and IBH Publishing Co. Pvt. Ltd., New Delhi.

#### Reference Books

- **1.** Wiley JM, Sherwood LM and Woolverton CJ. (2013) **Prescott's Microbiology**. 9<sup>th</sup> Edition. McGraw Hill International.
- **2.** Jacquelyn G. Black (2015). **Microbiology: Principles and Explorations.** 9<sup>th</sup> Edition. John Wiley and Sons Australia Limited.
- **3.** Kathleen Park Talaro (2014). **Foundations in Microbiology: Basic Principles**, 9<sup>th</sup> Edition. McGraw-Hill Higher Education.
- **4.** Tortora GJ, Funke BR and Case CL. (2016). **Microbiology: An Introduction**. 11<sup>th</sup> Edition. Pearson Education Limited.
- **5.** Madigan MT, Martinko JM, Dunlap PV and Clark DP. (2014). **Brock Biology of Microorganisms**. 14<sup>th</sup> edition. Pearson International Edition
- **6.** Atlas RM. (1997). **Principles of Microbiology**. 2<sup>nd</sup> edition. WM.T. Brown Publishers. Hill Book Company.
- **7.** Stanier RY, Ingraham JL, Wheelis ML, and Painter PR. (1999). **General Microbiology**. 5<sup>th</sup> edition. McMillan.

## **Web References**

- 1. https://www.britannica.com/science/microbiology
- 2. https://nptel.ac.in/courses/102103015/pdf/mod8.pdf
- 3. https://www.atsu.edu/faculty/chamberlain/Website/Lects/Content1.html

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓		✓	✓	✓
CO2	✓	✓	✓		✓
CO3	✓	✓	✓	✓	
CO4	✓		✓	✓	
CO5	✓	✓		✓	

(For the candidates admitted from 2018- 19 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

First Semester Microbiology

## PRINCIPLES OF MICROBIOLOGY

Time: Three hours Maximum Marks: 75

**PART - A**  $(20 \times 1 = 20 \text{ Marks})$ Answer **ALL** the Questions All questions carry equal marks

	An questions early equal marks.
1.	Chondroid of some bacteria are better known as
	a. Bacterial mitochondria b. Mesosomes c. Bacterial plastids d. Plasmids
2.	The resolving power of an optical microscope is
	a. 0.2µm b. 0.2 Å c. 0.2 nm d. 0.2 mm
3.	Which of the following structure is absent in Gram positive bacteria?
	a. Cell wall b. Teichoic acid c. Murein d. Outer membrane
4.	Bacterial cells can be stained withto reveal the presence of lipid inclusions
	a. Saffranin b. Methylene blue c. Trypan blue d. Sudan dyes
5.	Who discovered Mycobactyerium tuberculosis?
	a. Koch b. Jenner c. Pasteur d. Virchow
6.	Who discovered Bacillus anthracis?
	a. Koch b. Pasteur c. Jenner d. Hansen
7.	Scientist who discovered theory of spontaneous generation
	a. Koch b. Pasteur c. Jenner d. Hansen
8.	The iodine used in Gram staining serves as
	a. Chelator b. Catalyst c. Mordant d. Cofactor
9.	The organism which obtain their energy from chemicals are designated as
	a. Prototroph b. Chemotrophs c. Organotrophs d. Autotrophs
10	. In the process of freeze drying, a dense cell suspension is placed in small vials and is
	frozen at
	a60 to 78°C b20 to -30 °C c30 to -48 °C d48 to -58 °C
11.	. Which of the following may contain fimbriae
	a. G+ ve bacteria b. G-ve bacteria c. Both A and B d. None of these
12	. Which were the investigators lived at the same time?
	a. Koch and Pasteur b. Darwin and Woese
	c. Leeuwenhoek and Ricketts d. Berg and Hooke
13.	Which of the following articles can be sterilized in an autoclave?
	a. Gloves b. Culture media c. Dressing material d. All of these
14.	Which of the following is not a disinfectant containing a heavy metal?
	a. Silver nitrate b. Mercurochrome c. Copper sulphate d. Chlorine

- 15. The oldest eukaryotic organisms are consider to be
  - a. Diplomonads like Giardia
- b. Archaea
- c. Fungi
- d. Animals
- 16. Which of the following is considered the most unifying concept in biology?
  - a. Taxonomy b
- b. Anatomy
- c. Genetics
- d. Evolution
- 17. Which of the following structure is absent in eukaryotic cell?
  - a. Mitochondria
- b. Chloroplasts
- c. Golgi structure
- d. Mesosome

- 18. The five kingdom system of classification was set up by
  - a. Louis Pasteur
- b. Robert Whittaker
- c. Robert Koch
- d. Masaki Ogata
- 19. Which of the following bacteria lack a cell wall and are therefore resistant to penicillin?
  - a. Cyanobacteria
- b. Mycoplasma
- c. Bdellovibrios
- d. Spirochetes
- 20. Which of the following best represents the hierarchy of levels of biological classification?
  - a. Phylum, kingdom, class, order, genus, species, family
  - b. Kingdom, phylum, class, order, family, genus, species
  - c. Kingdom, phylum, family, class, order, genus, species
  - d. Class, order, kingdom, phylum, family, genus, species

## **PART – B** (5 x 5 = 25 Marks)

Answer **ALL** questions

All questions carry equal marks

- 21. (a) What are Koch's postulates (or)
  - (b) Write about the scope of microbiology.
- 22. (a) Write about dark field microscope (or)
  - (b) Write short notes on Gram staining.
- 23. (a) Write short notes on transport media (or)
  - (b) What are antibiotics? Write about their types.
- 24. (a) Write an account on numerical taxonomy (or)
  - (b) Write short notes on Whittaker's five kingdom classification.
- 25. (a) Give an account of thermophiles (or)
  - (b) Briefly explain about actinomycetes.

## **PART - C** $(3 \times 10 = 30 \text{ Marks})$

**Answer ANY THREE Questions** 

All questions carry equal marks

- 26. Write a brief account on the historical developments of microbiology.
- 27. Write about Phase contrast microscope and their applications in microbiology.
- 28. Write in detail about the physical methods of sterilization.
- 29. Give an account of classification of bacteria according to Bergey's manual of systematic bacteriology.
- 30. Give a brief account on microbial diversity on diverse environment.

## PRINCIPLES OF MICROBIOLOGY (PRACTICALS)

## **Objectives**

- To introduce the Good laboratory practices and biosafety
- To learn the SOP of basic instruments in microbiology lab
- To cultivate the microbes in laboratory
- To learn the basic techniques leading to characterization of microbes
- To evaluate the antibiotic sensitivity pattern of microbes

## **Course Outcome:**

CO1	The knowledge on microbiology laboratory, working practices, basic instruments to
	be imparted
CO2	The handling of microscope for visualizing the morphology, size and movement of
	microbes could be learnt
CO3	The non pathogenic microbial cultivation may be practiced
CO4	The enumeration techniques from various samples may be experienced
CO5	The efficacy of the antibiotic sensitivity test might be learnt

- 1. Microbiology Good Laboratory Practices and Biosafety.
- 2. The principle and applications of instruments (biological safety cabinets, autoclave, incubator, hot air oven, light microscope, pH meter) used in the microbiology laboratory.
- 3. Preparation of culture media for aerobic and anaerobic bacteria.
- 4. Pure culture technique- Serial dilution, pour plate, spread plate and streak plate.
- 5. Enumeration of bacteria and actinobacteria from environmental sample (soil/water).
- 6. Staining techniques- simple, differential, negative, endospore, capsular, metachromatic granules and flagellar staining.
- 7. Determination of bacterial motility by hanging drop technique.
- 8. Antibiotic sensitivity test by Kirby Bauer method.

## **Suggested Reading**

1. Cappucino J and Sherman N (2010). **Microbiology: A Laboratory Manual**. 9<sup>th</sup> edition. Pearson Education Limited.

- 2. P. Gunasekaran (2005). **Laboratory Manual in Microbiology**. 1<sup>st</sup> Edition. New Age International Publishers.
- 3. Mette Praetorius Ibbe and Katherine Elasky (2017). **Basic and Practical Microbiology Laboratory Manual**. 1<sup>st</sup> Edition. Cognella. Incorporated.
- 4. Norbel A. Tabo (2004). Laboratory Manual in Microbiology. 1st Edition. Rex Book Store.
- 5. N.Kannan (2002). **Laboratory Manual in General Microbiology**. 1<sup>st</sup> Edition. Panima Publishing Corporation.
- 6. Sundara Rajan. S (2001). **Practical Manual of Microbiology**. 1<sup>st</sup> Edition. Anmol Publication Private.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓		✓
CO2	✓	✓	✓	✓	
CO3			✓	✓	✓
CO4	✓	✓		✓	
CO5	✓	✓	✓		✓



CORE - II

**Total Number of Hours: 60** 

4 Hours/ Week

## MICROBIAL PHYSIOLOGY AND METABOLISM

## **Course Objectives:**

- To study the Cellular structure of prokaryotes and eukaryotes.
- To gain knowledge about bacterial growth.
- To understand the transport mechanism of the bacteria.
- To study the metabolism and its types.
- To gain knowledge on mechanism of photosynthesis in bacteria.

#### **Course Outcome:**

CO1	The difference between the Eukaryotic and Prokaryotic cellular organizations were
	understood
CO2	The student got a clear idea of the bacterial growth and the factors influencing the
	growth
CO3	The different methods involved in the transport of materials from outside
	environment into the bacterial cell were taught
CO4	The metabolism of microbes with reference to different cycles were learnt
CO5	The microbial respiration and its classification based on the respiration were studied

UNIT – I No. of Hours: 12

**Cellular structures of prokaryotes:** Ultra structure and Functions of Prokaryotic cell wall, flagella, slime layer, capsule, pili, cytoplasmic membrane and cytoplasmic inclusions – Structure and functions of cyanobacteria.

UNIT – II No. of Hours: 12

**Growth of bacteria:** Nutritional types of bacteria – nutritional requirements – factors influencing microbial growth – growth curve – Generation time - Mathematical determination of growth. Multiplication, Sporulation and its mechanism.

UNIT – III No. of Hours: 10

**Microbial nutrients and transport:** Nutrients – Synchronous, Batch, continuous and diauxic growth culture. Structure and organization of membrane – Methods of nutrients transport in bacteria – Diffusion, active transport and facilitated diffusion – group translocation.

UNIT – IV No. of Hours: 14

**Aerobic respiration and Fermentation:** glycolysis, pentose phosphate pathways, EMP, TCA and Glyoxalate cycle - ATP synthesis and utilization – photophosphorylation, oxidative phosphorylation, substrate level phosphorylation - Fermentation types – Lactic acid, Butanol and Propionic acid. Respiration types – aerobic and anaerobic respiration.

UNIT – V No. of Hours: 12

**Anaerobic respiration:** Characteristics and metabolism of autotrophs - autotrophic CO<sub>2</sub> fixation and mechanism of photosynthesis – Oxygenic (cyanobacteria) and Anoxygenic (purple sulfur, green sulfur and halobacteria) – Physiology of Bio luminescence, Nitrogen fixation.

#### **Text Books**

- 1. Pelczar MJ, Chan ECS and Kreig NR (2008). **Microbiology**. 5<sup>th</sup> Edition, Tata McGraw Hill-Hill Education Pvt. Ltd., New Delhi.
- 2. Ram Reddy S and Reddy SM (2005). **Microbial Physiology.** 1<sup>st</sup> Edition. Scientific Publishers, India.
- 3. Meenakumari S (2006). **Microbial Physiology**. 1<sup>st</sup> Edition.MJP Publishers, A unit of Tamil Nadu Book House, Chennai.

## **Reference Books**

- 1. Moat G, John W Foster and Michael P Spector (2002). **Microbial Physiology.** 4<sup>th</sup> Edition. Wiley-Lis, Inc., New York.
- 2. Daniel R. Caldwell (2000). **Microbial Physiology and Metabolism.** 2<sup>nd</sup> Edition. Star Publishing Company.
- 3. Willey, J.M., Sherwood, L and Wool Verton C.J. (2011). **Prescott's Microbiology.** 8<sup>th</sup> edition, McGraw Hill, New York.

## Web sources

- 1. https://nptel.ac.in/courses/122103039/pdf/mod4.pdf
- 2. https://nptel.ac.in/courses/102103015/19
- 3. https://www.cliffsnotes.com/study-guides/biology/biology/the-biology-of cells/prokaryote-and-eukaryote-cell-structure

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	<b>✓</b>	✓	✓	✓	✓
CO2	✓	✓	✓		✓
CO3	<b>✓</b>	✓	✓	✓	
CO4	✓		✓	✓	✓
CO5	✓	✓		✓	✓

(For the candidates admitted from 2017- 18 onwards)

#### **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Second Semester Microbiology

#### MICROBIAL PHYSIOLOGY AND METABOLISM

Time: Three Hours Maximum Mark: 75

PART - A (20 x 1 = 20 Marks)

Answer ALL questions

All questions carry equal marks

- 1. Bacterial cell wall is made up of
  - a. Chitin b. Cellulose c. Dextran
- d. Peptidoglycan

d. Spinin

- 2. Bacterial flagella is made up of
  - a. Microtubules
- b. Tubulin
- c. Flagellin
- 3. Surface appandages of bacteria on cell wall attachment during conjugation is
  - a. Pili b. Flagella
- c. Spinae
- d. Cilia
- 4. The region where bacterial genome resides is called as
  - a. Nucleus b. Cytoplasm c. Nucleiod d. Ribos
    - d. Ribosome free region
- 5. Bacteria reproduce vegetatively by
  - a. Fission only

- b. Fission and fragmentation
- c. Fission, fragmentation and budding
- d. None of the above
- 6. Growth in a closed system, affected by nutrient limitation and waste product accumulation is called as ----
  - a. Batch culturing b. Ascus c. Fruiting body d. Continuous culturing
- 7. The organisms that obtain energy from chemicals are called
  - a. Prototrophs b. O
    - b. Organotrophs
- c. Chemotrophs
- d. Autotrophs
- 8. Which of the following is the characteristics of a growth curve
  - a. shows development of microbial population under relatively stable environmental conditions
  - b. plotted with logarithmic numbers
  - c. graph numbers of microbes versus time
  - d. each growth curve consists of four distinct phases
- 9. The significance of plasma membrane is that
  - a. it selectively allow a some molecules to pass into the organism
  - b. it prevents movement of molecules out of the organism
  - c. it is the site of protein synthesis
  - d. All of the above
- 10. The most important role of the prokaryotic cell wall is to
  - a. maintain the shape of the cell wall
  - b. protect the cell from osmotic pressure
  - c. prevent ions from diffusing away from the cell
  - d. block the effects of antibiotics like penicillin
- 11. ----- protein combines with the substance and helps to move across the membrane
  - a. Carrier
- b. Channel
- c. Cell recognition
- d. Receptor

- 12. Which of the following describes the fluid mosaic model of the plasma membrane structure
  - a. Phospholipid monolayer with embedded proteins
  - b. Phospholipid bilayer with embedded proteins
  - c. Phospholipid trilayer with embedded proteins
  - d. Triglyceride bilayer with embedded proteins
- 13. Hetero lactic bacteria produce ----
  - a. lactic acid only

- b. lactic acid + water + carbon di oxide
- c. lactic acid + carbon di oxide
- d. lactic acid + alcohol + carbon di oxide
- 14. In aerobic respiration, the terminal electron acceptor is
  - a. Oxygen
- b. Nitrogen
- c. Hydrogen
- 15. The process of converting chemical energy into chemical bond of ATP is called ---
  - a. Glycolysis
- b. Conversion
- c. Cellular aspiration
- d. Energy
- 16. The light trapping pigment molecule in plant, algae and cyanobacteria
  - a. Chlorophyll a
- b. Chlorophyll b
- c. Porphyrin
- d. Rhodopsin
- 17. The oxygen released into the air as a product of photosynthesis comes from -----a. Chlorophyll b. Carbon di oxide c. Water d. None of the above
- 18. Which of the following does not produce oxygen as a product of photosynthesis
- - b. Purple Sulphur bacteria c. Cyanobacteria d. Phytoplankton a. Oak trees
- 19. Hexose monophosphate pathway is also known as
  - a. Phosphogluconate pathway
- b. Oxalocaetate pathway

c. Malate pathway

- d. Fumerate pathway
- 20. The glyoxylate cycle is used by some organisms when ---- is the sole carbon source
  - a. Acetate
- b. Nitrate
- c. Carbon di oxide
- d. All of the above

## $PART - B (5 \times 5 = 25 \text{ Marks})$

## Answer **ALL** questions

All questions carry equal marks

- 21. (a) Write about the cell wall structure of bacteria (or) (b) Write a short note on capsule.
- 22. (a) Add a brief account on growth curve (or)
  - (b) Write about the nutritional requirements of microbes.
- 23. (a) Explain the fluid mosaic model of cell membrane (or) (b) Describe passive diffusion.
- 24. (a) Explain Kreb's cycle (or) (b) Explain mixed acid fermentation.
- 25. (a) Briefly describe the metabolism of autotrophs (or)
  - (b) Write an account on anoxygenic photosyntheis.

## $PART - C (3 \times 10 = 30 \text{ Marks})$

## Answer **ANY THREE** questions

All questions carry equal marks

- 26. Explain in detail about the mechanism of sporulation.
- 27. Explain the various factors that affecting the microbial growth.
- 28. Describe the various mechanisms of active transport.
- 29. Discuss in detail about microbial photosynthesis.
- 30. Explain briefly about the Physiology of Biolumninescence.

Total Number of Hours: 45
3 Hours/ Week

## MAJOR PRACTICAL - II - MICROBIAL PHYSIOLOGY AND METABOLISM

## **Course Objectives**

- To study the bacterial growth
- To study the effect of temperature, pH, carbon, nitrogen and salt concentration, incubation time, inoculums size on bacterial growth
- To understand the characterization of unknown organisms

## **Course Outcome:**

CO1	Different stages of bacterial growth could be studied
CO2	The impact of different physical parameters on bacterial growth are to be learnt
CO3	The impact of different chemical parameters on bacterial growth are to be learnt
CO4	The characterization of microorganisms based on IMViC tests are to be introduced
CO5	The characterization of microorganisms based on sugar assimilation are to be
	introduced

- 1. Bacterial growth curve Turbidometric assay.
- 2. Effect of temperature on growth of bacteria.
- 3. Effect of pH on growth of bacteria.
- 4. Effect of carbon and nitrogen sources on growth of bacteria.
- 5. Effect of salt concentration on growth of bacteria.
- 6. Effect of incubation time and inoculum size on growth of bacteria
- 7. Biochemical parameters
  - a) IMViC
  - b) Sugar assimilation (glucose, lactose, maltose, mannitol and sucrose)
  - c) Catalase
  - d) Oxidase
  - e) Urease
  - f) TSI

## **Reference Books**

1. Cappucino J and Sherman N. (2010). **Microbiology: A Laboratory Manual**. 9<sup>th</sup> edition. Pearson Education Limited.

- 2. P.Gunasekaran. (2005). **Laboratory Manual in Microbiology**. 1<sup>st</sup> Edition. New Age International Publishers.
- 3. Mette Praetorius Ibbe and Katherine Elasky. (2017). **Basic and Practical Microbiology Laboratory Manual**. 1st Edition. Cognella. Incorporated.
- 4. Norbel A.Tabo. (2004). Laboratory Manual in Microbiology. 1st Edition. Rex Book Store.
- 5. N.Kannan. (2002). **Laboratory Manual in General Microbiology**. 1<sup>st</sup> Edition. Panima Publishing Corporation.
- 6. Sundara Rajan. S. (2001). **Practical Manual of Microbiology**. 1<sup>st</sup> Edition. Anmol Publication Private.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓		✓	✓	
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓

SEMESTER – II
18U2MBA01 Total number of Hours: 60
Credits: 4 4 Hours/Week

## **BIOINSTRUMENTATION TECHNIQUES**

## **Course Objectives:**

- To gain knowledge about laboratory requirement for microbiology laboratory
- To study the recent advancements in chromatography
- To impart knowledge on Electrophoretic techniques and its applications
- To study the different types of centrifuges
- To understand spectroscopic techniques

## **Course Outcome:**

CO1	The course emphasizes on the basics of laboratory, its requirements and rules. It also gives
	an understanding about the recent advancements in microscopy, principle and the
	operation of the basic equipments used in the microbiology/clinical laboratory
CO2	Provides basic principles and separation of molecules by various chromatography
	techniques
CO3	Able to uptake introductory principle and background of electrophoresis and its common
	application in separation of genetic material and high throughput techniques for the
	separation of biomolecules
CO4	It is an opportunity to understand the working principles of analytical spectrophotometers
	and its applications
CO5	Ability to understand the most common and routine laboratory separation of molecules based on physical and chemical properties

UNIT – I No. of Hours: 10

**Microbiological Instruments:** Basic requirements of a Microbiology Laboratory – Basic microbiological Instruments - Colony counter, Neubaeur chamber, inoculation loop, transilumintor, cyclo mixer, Incubator - shaker incubator, BOD incubator, CO<sub>2</sub> Incubator. Microscopy – SEM, TEM and confocal. Balance types – mono pan, top and physical.

UNIT – II No. of Hours: 12

**Centrifugation:** Centrifuge – Sedimentation principle, Relative centrifugal force, Sedimentation coefficient, factors affecting sedimentation velocity, Centrifuge rotors, Types of centrifuges, Ultracentrifuge – Preparative and Analytical – Centrifugation – Types - Differential and Density gradient centrifugation.

UNIT – III No. of Hours: 10

**Spectrophotometry:** Principle and applications – Beer Lambert's Law – Principle and applications of Colorimeter, spectrophotometer, Atomic Adsorption Spectrophotometer, Raman spectrophotometer – Analysis of biomolecules using UV and visible spectrophotometer – Spectroflourimeter.

UNIT – IV No. of Hours: 14

**Electrophoresis:** Principle and applications – Agarose gel electrophoresis, Pulse Field Gel Electrophoresis, SDS – polyacrylamide gel electrophoresis, 2D gel electrophoresis, Isoelectric focusing and Zymogram preparation.

UNIT – V No. of Hours: 14

**Chromatography:** Principles and applications of paper chromatography (including Descending and 2-D), Thin layer chromatography, Gel filtration chromatography, Ion-exchange chromatography, affinity chromatography, HPLC and GC-MS.

#### Text Books

- 1. Praful K Godkarand and Darshan P Godkar (2006). **Text book of Medical Laboratory Technology.** Bhalani Publishing House, Mumbai.
- 2. Arora CK and Prakash M (1998). **Laboratory instrumentation.** Anmol Publications Pvt. Ltd., New Delhi.

## **Reference Books**

- 1. Keith Wilson and John Walker (1994). **Principles and Techniques of Practical Biochemistry.** 5<sup>th</sup> Edition, Cambridge University Press, New York.
- 2. Rodney Boyer (2000). **Modern Experimental Biochemistry.** 3<sup>rd</sup> Edition, Addition Wesley Longman, San Francisco.
- **3.** Webster JG (2004). **Bioinstrumentation**. University of Wisconsin, John Wiley & Sons, Inc. UK.

## **Web Sources**

- 1. https://nptel.ac.in/courses/102103044/3
- 2. http://www.biologydiscussion.com/microbiology-2/instruments/top-9-instruments-used-in-microbiology/66905

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	
CO2	<b>✓</b>		✓	✓	<b>✓</b>
CO3		✓	✓		✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Second Semester Microbiology

## **BIOINSTRUMENTATION TECHNIQUES**

Tim	e: Three Hours			Maximum Mark:	<b>75</b>
		PART	- <b>A</b> (20 x 1= 20 N	Marks)	
		An	swer ALL questio	ons	
		All que	estions carry equal	marks	
	1. The instrument	used for homogenous	mixing is		
	a. Incubator	b. Cyclomixer	c. Centrifuge	d. Shaker	
	2. The modern ver	rsion of counting colon	y is		
	a. Digital Coun	ter b. Colony Counte	er c. Ultra Cou	unter d. Mechanical Counter	
	3. The source of la	ight used in transillumi	nator is		
	a. Sodium vapo		c. Fluorescent	d. IR	
	4. Who first descr	ibed colony counter	• • • • • • • • • • • • • • • • • • • •		
	a. Robert Koch	•		d. Antony von Leewenhoek	
	•	e for imaging in electro	n microscope		
	a. Neutron	b. Electron	c. Proton	d. All the above	
		n technique used for inc	-		
	a. alcohol	b. incineration	c. boiling	d. Pasteurization	
	•	emperature for bacteria	· ·		
	a. 35°C	b. 36°C	c. 37°C	d. 38°C	
		rmally used in microbio			
	a. Top pan	b. Mono pan	c. Physical	d. Chemical	
		by in Agarose	-		
	a. EtBr	b. Bromothymol b		d. Bromophenol blue	
		nnique used for the sepa	_		
	a. RNA	b. DNA	c. Protein	d. Lipid	
	11. PAGE stands		1 D 1	1 11 11 11	
	• •	ide gel electrophoresis	• •	lamide gel electrophoresis	
		el electrophoresis	, ,	l electrophoresis	
		s an electrophoretic tec	-		
	• •	zymes b. Proteins	•	d. substrates	
		b. semiquatitative		analysis of molecules	
	a. quantative	•	•	•	
	14. A spectrophot	ometer is an instrumen	t that measures the	e amount of photons	

b. semiquatitative c. qualitative

d. semiqualitative

a. quantative

15. Beer-Lambert Law states that there is a linear relationship between the absorbance and the --------- of a sample. a. concentration b. strength d. equivalence c. amount 16. The proportion of light absorbed by a medium is ----- of the intensity of incident light. a. independent b. dependent c. direct d. indirect 16. RPM means. a. rotation per minute b. reel per minute c. random per minute d. redeem per minute 17. Density gradient centrifugation Is considered one of the more efficient methods of a. separating suspended particle b. separating particle c. suspended particle d. particle 18. The forces involved in centrifugation is c. gravity a. Centripetal b. centrifugal d. external 19. CeCl is a type of a. gradient centrifuge b. Normal centrifuge c. differential centrifuge d. microfuge 20. The shaft is attached with b. motor d. bucket a. rotor c. rod

## **PART - B** (5 x 5 = 25 Marks)

Answer **ALL** questions

All questions carry equal marks

- 21. (a) Write about structure and application of BOD incubator (OR)
  - (b) Write a short note on transiluminator.
- 22. (a) Add a brief account on paper chromatography (OR)
  - (b) Describe about thin layer chromatography.
- 23. (a) Explain about the principle and application of agarose gel electrophoresis (OR)
  - (b) Describe about Zymogram preparation.
- 24. (a) Explain Beer Lambert's law (OR)
  - (b) Explain about spectroflourimeter.
- 25. (a) Briefly describe the sedimentation principle (OR)
  - (b) Write an account on density gradient centrifugation.

## **PART** – **C** (3 X 10 = 30 Marks)

Answer **ANY THREE** questions

All questions carry equal marks

- 26. Explain in detail about the mechanism and application of electron microscope.
- 27. Explain the various factors that affecting chromatography.
- 28. Describe in detail about SDS electrophoresis.
- 29. Discuss in detail about UV Spectrophotometer.
- 30. Explain briefly about the ultracentrifuge.

## ALLIED PRACTICAL – II - BIOINSTRUMENTATION TECHNIQUES

## **Course Objectives:**

- To know about the basics of solution preparation for various experiments
- To get trained in the estimation of biomolecules
- To understand the working principle and methods of analytical instruments
- To get skilled in basic molecular biology techniques
- To get trained in basics of chromatography

#### **Course Outcome:**

CO1	Become well-versed in preparation of reagents and buffers				
CO2	It offers to participate with very advanced chromatographic methods for the separation				
	of molecules				
CO3	The student can learn most common methods to separate genetic material and proteins				
CO4	Allows to capture detailed working principle of spectrophotometry and its application				
CO5	A hands on approach to develop skill in estimation of biomolecules using spectrophotometry				

- 1. Calculation in preparation of reagents: Normality of solution, Molarity of solution
- 2. Chromatographic Techniques: Paper and Thin layer chromatography
- 3. **Electrophoretic Techniques:** Agarose gel electrophoresis, SDS-PAGE
- 4. **Spectrophotometry**: Principle and operating mechanism of Spectrophotometry, Estimation of biomolecules like Protein and Carbohydrate and Lipid using UV and visible Spectrophotometer.

## **Reference Books**

- 1. Rodney Boyer (2000). **Modern Experimental Biochemistry.** 3<sup>rd</sup> Edition, Addition Wesley Longman, San Francisco.
- 2. John G Webster (2004). **Bioinstrumentation**. University of Wisconsin, John Wiley & Sons, Inc. U K.
- 3. Keith Wilson and John Walker (1994). **Principles and Techniques of Practical Biochemistry**. 5<sup>th</sup> Edition, Cambridge University Press, New York.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓



SEMESTER – III

18U3MBC03

Credits – 4

Core – III

Total number of Hours: 60

4 Hours/Week

## MOLECULAR BIOLOGY AND MICROBIAL GENETICS

## **Course Objectives:**

- To gain knowledge about DNA and RNA
- To understand DNA replication and transcription in prokaryotes & eukaryotes
- To impart knowledge on translation and gene regulation
- To study the features of plasmid and mechanism of genetic exchange
- To gain knowledge about mutation and repair mechanisms

#### **Course Outcome:**

CO1	It enables to understand the historical perspective and background / basic knowledge of
	genetics
CO2	It gives exposure on central dogma of life
CO3	It helps to uptake knowledge on translation and gene regulation in prokaryotes
CO4	It delivers basic knowledge and techniques used in gene transfer
CO5	It provides basic concepts of mutation and mutagenesis and gene repair mechanisms

UNIT – I No. of Hours: 12

**Genetic Material (DNA & RNA):** Genetics – Historical perspectives, discovery of DNA structure – Watson and Crick model – Types and forms of DNA, Genome organization in Prokaryotes, Viruses and Eukaryotes. DNA as a genetic material. RNA as genetic material. Physical structure and chemical composition of nucleic acid. RNA types – t RNA, mRNA and rRNA.

UNIT – II No. of Hours: 12

**Replication and transcription:** DNA replication in prokaryotes – Meselson and Stahl experiment – Mechanism and enzymology of replication - Rolling circle and theta model of replication. Transcription in prokaryotes: promoter, RNA polymerase.

UNIT – III No. of Hours: 12

**Translation:** Salient features of genetic code - Wobble hypothesis. Translational machinery, charging of tRNA, aminoacyl tRNA synthetases, Mechanisms of initiation, elongation and termination in prokaryotes. Operon concept – *lac* and *trp* operons.

UNIT – IV No. of Hours: 12

**Plasmid:** Types of plasmids – F plasmid, R Plasmids, Col plasmids, Ti plasmids, Plasmid replication and partitioning, Host range, plasmid –Incompatibility, plasmid amplification, Regulation of copy number, curing of plasmids.

UNIT – V No. of Hours: 12

Gene transfer, Mutation and DNA repair mechanisms: Transformation – Discovery, mechanism of natural competence. Conjugation – mechanism, Hfr and F' strains, Transduction – Generalized transduction, specialized transduction. Mutations and types of mutation - Auxotrophic mutant detection: Replica plate technique. Mutagenicity testing – Ames Test. DNA repair mechanisms – excision, mismatch, SOS, photoreactivation and recombination repair.

#### **Text Books**

- 1. David Freifelder (2005). **Molecular Biology**. 2<sup>nd</sup> Edition. Narosa Publishers, New Delhi.
- 2. Verma PS and Agarwal VK (2006). **Cell Biology, Genetics, Molecular Biology, Evolution and Ecology.** S. Chand & Company Ltd., New Delhi.

#### Reference Books

- 1. Friedberg EC, Walker GC, Siede W (2006). **DNA repair and mutagenesis**. ASM press, Washington DC.
- 2. Benjamin Lewin (2000). Genes VII. 7th Edition. Oxford University press, Inc.
- 3. Maloy SR, Cronan JE, FreifelderD (1994). Microbial Genetics. Jones and Bartlett Publishers.
- 4. Gardner EJ, Simmons MJ, Snustad DP (2008). Principles of Genetics. 8th Ed. Wiley-India.
- 5. Watson JD, Baker TA, Bell SP, Gann A, Levine M and Losick R (2008). **Molecular Biology** of the Gene, 6<sup>th</sup> edition, Cold Spring Harbour Lab. Press, Pearson Publication

#### Web sources:

- 1. http://biology.kenyon.edu/courses/biol63/watson 06.pdf
- 2. https://nptel.ac.in/courses/102103015/33
- 3. https://nptel.ac.in/courses/102103017/module26/lec26\_slide9.htm

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	<b>✓</b>			✓	
CO2	✓		✓	✓	✓
CO3		✓	✓		✓
CO4	✓	✓		✓	
CO5	✓	✓	✓		✓

(For the candidates admitted from 2017- 18 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Third Semester Microbiology

MOLECULAR BIOLOGY AND MICROBIAL GENETICS
Time: Three Hours Maximum Mark: 75
PART - A (20  x  1 = 20  Marks)
Answer ALL questions
•
All questions carry equal marks
1. A nucleoside is composed of
a) a base+ a sugar b) a base+ a sugar+ phosphate c) a base+ a phosphate d) none of these
2. Two strands in a DNA double helix is joined by
a) Covalent bond b) Hydrogen bond c) Ionic bond d) Phosphodiester bond
3. The sugar in RNA Is
a) Deoxyribo sugar b) Ribo sugar c) Fructose d) Sucrose
4. Thymine in DNA replaced by
a) Uracil b) Adenine c) Guanine d) Cytosine
5. Which enzymes remove supercoiling in replicating DNA ahead of the replication fork?
a) Helicases b) DNA polymerases c) Primases d) Topoisomerases
6. During which phase of the cell cycle is DNA replicated?
a) G1 phase b) S phase c) G2 phase d) M phase
7. True replication of DNA is possible due to
a) Hydrogen bonding b) Phosphate backbone c) Complementary base pairing d) None of above
8. Most of mistakes during DNA replication are corrected by
a) DNA polymerase b) DNA ligase c) gyrase d) Helicase
9. The accepted hypothesis for DNA replication is
a) Conservative theory b) Dispersive theory c) Semi-conservative theory d) Evolutionary theory
10. Telomeres are usually rich in which nucleotide?
a) Adenine b) Guanine c) Cytosine d) Thymine
11. In prokaryotes, the first aminoacid in the polypeptide chain is
a) Methionine b) N-Methyl methionine c) N-Formyl methionine d) N-Acetyl methionine
12. On which of the following molecules would you find a codon?
a) mRNA b) rRNA c) tRNA d) SnRNA
13. What amino acid is coded by the triplet of bases UAU?
a) Phenylalanine b) Tyrosine c) Serine d) Cysteine
14. Ti plasmid that is used as a plant vector is obtained from
a) Agrobacterium tumefaciens b) Agrobacterium rhizhogenes
c) Agrobacterium radiobacter d) Thermas aquaticus
15. Which of the following cells of <i>E.coli</i> are referred to as F-
a) Male b) Female c) Both A&B d) Neither A or B
16. Point mutation involves
a) Deletion b) Insertion c) Duplication d) Change in single base pair
17. When viral genome can become integrated into the bacterial genome they are known as
a) Temperate phage b) Prophage c) Bacteriophage d) Episome
18. Name a type of Radiation in induced mutations

- a) Microwave b) UV radiation c) Heat d) Both A & C
- 19. The function of enzyme involved in base excision repair is
- a) Addition of correct base
- b) Addition of correct nucleotide
- c) Removal of incorrect base
- d) Removal of phosphodiester bond
- 20. The enzyme photolyase is used in what method of repair?
- a) Base exicision
- b) Photoreactivation
- c) Nucleotide excision
- d) SOS repair

**PART - B** (5 x 5 = 25 Marks)

Answer **ALL** questions
All questions carry equal marks

- 21. Describe the Watson and Crick model of DNA (**OR**)
  - b) Discuss about the types of RNA.
- 22. a) Write short notes on Rolling circle model of replication (OR)
  - b) Explain about the bidirectional replication.
- 23. a) Write short notes on Lac operon (**OR**)
  - b) Differentiate the Transcription and Translation process.
- 24. a) Discuss about the Ti plasmid (**OR**)
  - b) Give short notes on Conjugation.
- 25. a) Write short notes on mutation and its types (**OR**)
  - b) Discuss about the Replica plate technique.

**PART – C** (3 X 10 = 30 Marks)

Answer **ANY THREE** questions

All questions carry equal marks

- 26. Briefly explain about the DNA as the genetic material.
- 27. Explain about the replication in prokaryotes.
- 28. Discuss the operon concept in detail.
- 29. Give a brief account on Transduction.
- 30. Explain briefly about the DNA Repair mechanisms.

CORE PRACTICAL – III
Total number of Hours: 30
3 Hours/Week

## MOLECULAR BIOLOGY AND MICROBIAL GENETICS (PRACTICALS)

## **Course Objectives:**

- i) To be aware of the isolation of chromosomal and plasmid DNA
- ii) To obtain knowledge on physical and chemical mutagenesis
- iii) To achieve knowledge about coli phage transfer method
- iv) To know about the gene transfer methods
- v) To get information about the techniques used in genetics

## **Course Outcome:**

CO1	The students would be skilled in chromosomal and plasmid DNA isolation from eukaryotes
CO2	They would be expertise with effects of physical and chemical agents responsible for mutagenesis
CO3	They can able to isolate antibiotic resistant and auxotrophic mutants
CO4	They would be exposed to hands on technique for the isolation of phage from sewage
CO5	They were enabled with fundamental techniques used for prokaryotic gene transfer techniques

- 1. Isolation of chromosomal DNA from bacteria
- 2. Isolation of plasmid DNA from *E. coli*
- 3. Physical and Chemical mutagenesis
- 4. Isolation of antibiotic resistant mutant by gradient plate technique
- 5. Isolation of auxotrophic mutant (replica plating)
- 6. Isolation of coli phage from sewage
- 7. Bacterial Gene Transfer Transformation (Demonstration)

#### Reference Books

- 1. Sambrook J and Russell DW (2001). **Molecular Cloning A laboratory manual.** 3<sup>rd</sup> Edition. Cold Spring Laboratory Press, New York.
- 2. Dubey RC and Maheshwari DK (2002). **Practical Microbiology**. S Chand and Co. Ltd., New Delhi.

- 3. Aneja KR (2010). **Experiments in Microbiology, Plant Pathology and Biotechnology.** New Age International (P) Limited Publishers.
- 4. Harold J Benson (2002). **Microbiological Applications: Laboratory manual in General Microbiology**. 8<sup>th</sup> Edition. Mcgraw-Hill, Boston.
- 5. James G Cappuccino and Natalie Sherman (2005). **Microbiology: A Laboratory manual.** 7<sup>th</sup> Edition, Pearson Education, Inc.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓			✓
CO2		✓		✓	✓
CO3		✓	✓	✓	✓
CO4	✓	✓		✓	✓
CO5	✓	✓	✓	✓	✓

SEMESTER – III

18U3MBA02

Credits – 4

ALLIED – III

Total number of Hours: 45

4 Hours/Week

## **BIOINFORMATICS**

# **Course Objectives:**

- 1. To gain knowledge about the fundamentals of computer.
- 2. To impart knowledge on bioinformatics and biological database.
- 3. To gain knowledge about sequence alignments and phylogeny tree.
- 4. To impart knowledge on genome organization.
- 5. To understand about protein structure prediction

#### **Course Outcome:**

CO1	It provides a full-fledged introductory knowledge on various databases used in
	bioinformatics
CO2	It offers direction for individual molecular databases and various tools used
CO3	It delivers a complete platform for the tools used for sequence alignment and
	phylogenetic tree construction
CO4	It provides a broad knowledge on drug designing and development and also
	phylogenetic tree construction
CO5	It imparts knowledge on the prediction of protein structures using various tools

UNIT – I No. of Hours: 09

**Introduction and Databases:** Introduction, database model, types of database - primary, secondary database, raw database and processed database, data mining, data storage and retrieval, querying in database and tools for querying – BLAST, FASTA.

UNIT – II No. of Hours: 09

**Introduction to Bioinformatics and Biological Databases:** Biological databases - nucleic acid, genome, protein sequence and structure, gene expression databases, Database of metabolic pathways, Mode of data storage - File formats - FASTA, Genbank and Uniprot, Data submission & retrieval from NCBI and PDB.

UNIT – III No. of Hours: 09

**Sequence Alignments:** Local and Global Sequence alignment, pairwise and multiple sequence alignment. Scoring an alignment, scoring matrices, PAM & BLOSUM series of matrices.

UNIT – IV No. of Hours: 09

**Drug designing and development, Phylogeny and Phylogenetic trees:** Druglikeness – ADME – toxicity – Docking studies. Types of phylogenetic trees, Different approaches of phylogenetic tree construction - UPGMA, Neighbour joining, Maximum Parsomony, Maximum likelihood.

UNIT – V No. of Hours: 09

**Protein Structure Predictions:** Hierarchy of protein structure - primary, secondary and tertiary structures, modeling. Structural Classes, Motifs, Folds and Domains. Protein structure prediction in presence and absence of structure template. Energy minimizations and evaluation by Ramachandran plot.

# **Suggested Reading**

- 1. Saxena Sanjay (2003). A First Course in Computers, Vikas Publishing House.
- 2. Pradeep and Sinha Preeti (2007). **Foundations of Computing**, 4<sup>th</sup> ed., BPB Publications.
- 3. Lesk M.A. (2008). **Introduction to Bioinformatics**. Oxford Publication, 3<sup>rd</sup> International Student Edition.
- 4. Rastogi S.C., Mendiratta N. and Rastogi P. (2007). **Bioinformatics: methods and applications, genomics, proteomics and drug discovery**, 2<sup>nd</sup> ed. Prentice Hall India Publication.
- 5. Primrose and Twyman (2003). **Principles of Genome Analysis & Genomics**. Blackwell.

## Web sources

https://nptel.ac.in/courses/102106065/14

https://nptel.ac.in/courses/102103044/39

http://www.cs.ubc.ca/labs/beta/Courses/CPSC536A-01/Class7/class7-notes.html

https://nptel.ac.in/courses/102103044/pdf/mod6.pdf

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓		✓	✓	
CO2	✓	✓	✓		✓
CO3		✓		✓	<b>✓</b>
CO4	✓	✓	✓		✓
CO5	✓		✓	✓	

(For the candidates admitted from 2017- 18 onwards)

# **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Third Semester Microbiology

# BIOINFORMATICS

	DI		
<b>Time: Three Hours</b>			Maximum Mark: 75
	PART -	- <b>A</b> (20 x 1= 20 Marks)	)
	Ans	wer ALL questions	
	All ques	tions carry equal mark	s
1. The first bioinfor	matics database is crea	ated by	
a. Richard durbin	b. Dayhoff c.	Micheal J.Dunn	d. PEarson
2. Operating system	n is		
a. Collection of har	dware components	b. Collection of Inpo	ut and output device
c. Collection of soft	tware routines	d. All of the baove	
3. The term bioinfo	rmatics was coined by		
a. JD Watson	b. Margaret dayhoff	c. Pauline hogweg	d. Freidric sanger
4. Stepwise method	in solving the problem	n in computer science is	s called
a. Flowchart	b. Sequential design		d. Algorithm
5. An example of H	lomology & similarity	tool	
a. PROSPECT	b. EMBOSS	c. RASMOL d. BI	LAST
6. Which of the foll	owing is nucleotide see	quence database	
a. EMBL	b. SWISSPROT	c. PROSITE	d. TrEMBL
7. SWISSPROT is	related to		
a. Portable data	b. Swiss bank data	c. Sequence data ba	nk d. Genome databank
8. Blast programme	e is used in		
	-	•	coding d. Bioinformaics
•	contains more than tw	•	
· · · ·	•	c. Global d. Lo	
• •	· ·	s called asal	· ·
a. global b. lo	1 0	d. non-progr	ressive
11. BLOSSUM mar			
• •	•	irwise sequence alignn	nent
c. Phylogenetic ana	•	ll the above	
	lationship can be show	•	
a. Dendrogram	b. Genebank c. Da		a. Data scaren toor
13. Which one of the	ne following is not a ch	aracter-based method i	n tree construction?

b. Minimum likelihood

d. Neighbor joining

a. Maximum parsimony

c. Minimum evolution method

- 14. Analysing or comparing entire genome of the species is called
- a. Bioinformatics
- b. Genomics
- c. Proteomics
- d. Pharmacogenomics

- 15. Human genome contains about -----
- a. 2 billion base pairs
- b. 3 billion base pairs
- c. 4 billion base pairs
- d. 5 billion base pairs

- 16. Two dimensional gels are used to separate
- a. DNA fragments
- b. RNA fragments
- c. different proteins
- d. DNA from RNA

- 17. The tool for identification of motif -----
- a. COPIA
- b. Patternhunter

19. A compound that have desirable properties of drug is called

- c. PROSPECT
- d. BLAST

- 18. Proteomic sis the study of
- a. Set of proteins

- b. Set of proteins in the specific region of cell
- c. Entire set of expressed proteins in cell
- d. none of these

- a. lead
- b. find
- c. fit drug
- d. fit compound
- 20. Hydropathy plots are usually used to predict -----
- a. beta secondary structure
- b. transmembrane domains
- c. alpha secondary structure d. tertiary structure

# **PART - B** (5 x 5 = 25 Marks)

Answer **ALL** questions

All questions carry equal marks

- 21. a) Write about data mining (**OR**)
  - b) Write a short note on raw and processed database.
- 22. a) Add a brief account on Protein sequence & structure database (**OR**)
  - b) Describe about Data submission in Primary database.
- 23. a) Explain about the local sequence alignment (**OR**)
  - b) Describe about Scoring in alignment with blosum matrics.
- 24. a) Explain about MALDI TOFF spectroscopy (**OR**)
  - b) Explain about the features of *E.coli* genome database.
- 25. a) Briefly describe the proteins motifs, folds & domains (**OR**)
  - b) Write an account on Ramachandran plot.

## $PART - C (3 \times 10 = 30 \text{ Marks})$

Answer **ANY THREE** questions

- 26. Explain in detail about the Primary & Secondary database.
- 27. Explain in detail about the nucleic acid database.
- 28. Describe in detail about Phylogenetic tree.
- 29. Discuss in detail about completed human genome database.
- 30. Explain briefly about the structure of protein.

## **BIOINFORMATICS (PRACTICAL)**

# **Course Objectives:**

- 1. To be familiar with sequence analysis
- 2. To be aware of docking studies
- 3. To achieve knowledge about phylogenetic analysis
- 4. To obtain knowledge about drug discovery and development
- 5. To get information about bioinformatics tools

## **Course Outcome:**

CO1	It provides technical information on LINUX operating system
CO2	It paves a way to study the essential bioinformatics databases and its analysis
CO3	It offers extensive knowledge on tools used for sequence alignment and phylogenetic analysis
CO4	It provides a broad knowledge on drug designing and development and docking studies
CO5	A comprehensive tools to analyze the proteins for their properties

- 1. Introduction to LINUX OS
- 2. Introduction to bioinformatics databases PUBCHEM/NCBI/PDB
- 3. Sequence analysis using BLAST FASTA
- 4. Phylogenetic analysis Neighbour joining tree
- 5. Drug discovery and development Druglikeness ADME toxicity
- 6. Docking studies

# **Suggested Reading**

- 1. Saxena Sanjay (2003). A First Course in Computers, Vikas Publishing House.
- 2. Pradeep and Sinha Preeti (2007). **Foundations of Computing**, 4<sup>th</sup> ed., BPB Publications.
- 3. Lesk M.A. (2008). **Introduction to Bioinformatics**. Oxford Publication, 3<sup>rd</sup> International Student Edition.

- 4. Rastogi S.C., Mendiratta N. and Rastogi P. (2007). **Bioinformatics: methods and applications, genomics, proteomics and drug discovery**, 2<sup>nd</sup> ed. Prentice Hall India Publication.
- 5. Primrose and Twyman (2003). Principles of Genome Analysis & Genomics. Blackwell.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	
CO2			✓		
CO3		✓	✓	✓	✓
CO4	✓	✓	✓		
CO5	✓			✓	✓

ALLIED – III Total Number of Hours: 45

4 Hours/ Week

## **MICROBIOLOGY**

# **Course Objectives:**

- To study the history of microbiology and to gain knowledge on microscopy
- To impart knowledge on bacterial anatomy and staining techniques
- To study the types of culture media, to understand sterilization techniques and to cultivate the microbes
- To understand the role of microbes in the field of medical, food and Environment

## **Course Outcome:**

CO1	Able to learn about chronological development and growth of microbiology and its
	importance and enables students to get motivated
CO2	It makes expertise in the art of techniques for the identification of microbes by
	staining methods
CO3	Enables to gather basic components of nutritional media, preparation and routine
	techniques used for the cultivation of microorganism in sterile condition
CO4	From this, one can stuff with medically and most prevalent diseases and its control /
	treatment
CO5	Helps to gain essential soil microbes and their significant role in agricultural field and
	food industry

UNIT – I No. of Hours: 09

**History & Scope of Microbiology**: Introduction - Contributions of various scientists to Microbiology - Louis Pasteur, Antony Van Leeuwenhoek, Robert Koch, Joseph Lister, Edward Jenner, Alexander Fleming. **Microscopy:** Bright field microscope, Dark field microscope, Phase contrast microscope, Fluorescent microscope and Electron microscope – TEM & SEM.

UNIT – II No. of Hours: 09

**Identification of Microbes**: Basic Structure of Bacteria – Gram positive and Gram negative bacteria. Stains and staining procedure - Types of staining - simple, differential and special staining – Fungal staining techniques – Lactophenol cotton blue staining and KOH mount.

UNIT – III No. of Hours: 09

**Cultivation of Microbes**: Culture media – Definition – Types - composition – Media preparation – Basal, Differential, Selective, Transport and Anaerobic culture media. Sterilization – Definition – Methods - Types of agents - Physical agents - Chemical agents. Culture techniques – Methods - Streak plate, Pour plate, Spread plate. Cultivation of anaerobes – Preservation of cultures.

UNIT – IV No. of Hours: 09

**Medical Microbiology:** Infection – Definition – Types – Mode of disease transmission – sources, Factors influencing pathogenesis – Disease cycle, Control of disease and prophylaxis. Peptic ulcer, Typhoid, Dengue, SARS, Candidiasis, Aspergillosis, Giardiasis.

UNIT – V No. of Hours: 09

**Applications of Microbiology:** Biofertilizer — Mycorrhiza, PGPR — Bioremediation — Biopesticides — Bacteria and Fungi, Biogas production - Bioactive compounds — Probiotics and prebiotics.

#### **Text Books**

- 1. Pelczar MJ, Chan ECS and Kreig NR (2008). **Microbiology**. 5<sup>th</sup> Edition, Tata McGraw Hill-Hill Education Pvt. Ltd., New Delhi.
- 2. Dubey RC and Maheswari DK (2005). **A Textbook of Microbiology**, Revised Multicolour Edition. S Chand and Company Limited, New Delhi.
- 3. Sullia S.B and Santhanam S (2005). **General Microbiology**. 2<sup>nd</sup> Edition, Oxford and IBH Publishing Co. Pvt. Ltd., New Delhi.

#### Reference Books

- 1. Kathleen Park Talaro (2009). **Foundations in Microbiology: Basic Principles**, 7<sup>th</sup> Edition. McGraw-Hill Higher Education
- 2. Stanier RY, Ingraham JL, Wheelis ML and Painter PR (1987). **General Microbiology**. 5<sup>th</sup> Edition, MacMillan Education Ltd., London.
- 3. Gerard J Tortora, Berdell R Funke, Christine L Case (2010). **Microbiology: An Introduction.** 10<sup>th</sup> Edition, Pearson Benjamin-Cummings Publishing Company.

## **Web References**

https://www.britannica.com/science/microbiology

https://www.atsu.edu/faculty/chamberlain/Website/Lects/Content1.htm

http://www.amm-mcrc.org/publications/Biofertilizers.pdf

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2			✓	✓	✓
CO3		✓	✓	✓	
CO4	✓	✓	✓		✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

# **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Second Semester Microbiology

ALLIED MICROBIOLOGY	
Time: Three Hours Maximum Mark: 75	
PART - A (20  x  1 = 20  Marks)	
Answer <b>ALL</b> questions	
All questions carry equal marks	
1. Robert Koch discovered	
a. Bacillus anthracis b. Salmonella typhi c. Ebola virus d. Amoeba parasite	
2. Anton Von Leeuwenhoek discovered	
a. Animalcules b. Virus c. Fungi d. Yeast	
3. Which one of the following is used to visualize live cells?	
a. Bright field microscopy  b. Dark field microscopy	
c. Phase contrast microscopy d. SEM	
4. Electron microscope was made by	
a. Robert hooke b. Knoll and Ruska c. Kepler and Galileo d. F.Janssen and Z.janssen	ì
5. Gram staining is the example for	
a. Simple staining b. Differential staining c. Special staining d. None of the above	
6. Lipopolysaccharide is found in cell wall of	
a. Gram positive bacteria b. Gram negative bacteria c. Both d. Fungi	
7. Which one of the following is used as disinfectant in LCB staining?	
a. Lactic acid b. Phenol c. Glycerol d. Methylene blue	
8. Which of the staining technique helps in demonstrating spore structure in bacteria as well as fr	ee
spores?	
a. Acid-fast stain b. Endospore stain c. Capsule stain d. Flagella stain	
9. Which of the following is a rich source of nitrogen?	
a. Peptone b. Yeast extract c. Beef extract d. Agar	
10. The importance of agar in culture media were discovered by	
a. Ehrlich b. Petri c. Finly d. Hessy	
11. During preservation of microbes their	
a. characteristics change b. metabolism stop c. metabolism continue d. metabolism change	
12. Which of the following method is widely used for the preservation of microbes?	
a. Drying in vacuum b. Storage in sterile soil	
c. Lyophilization d. Storage in saline	
13. Transmission of 'pathogens' during pregnancy from mother to child is called as	
<ul><li>a. Direct transmission</li><li>b. Horizontal transmission</li><li>c. Vertical transmission</li><li>d. Indirect transmission</li></ul>	
14. An insect or animal carrier of disease is known as	

c. Fomite

d. Vehicle

b. Vector

a. Carrier

- 15. Water quality is measured by
- a. MBRT
- b. Resazurin test
- c. Staining
- d. MPN

- 16. Cholera is caused by
- a. Vibrio
- b. E.coli
- c. Salmonella
- d. Pseudomonas

- 17. Pasteurization technique is used for
- a. Milk
- b. Cheese
- c. Bread
- d. Antiseptic
- 18. The undesirable change that makes the unsafe food consumption is called -----
- a. Food decay
- b. Food spoilage
- c. Food loss
- d. All the above

- 19. Botulism is caused by
- a. E.coli
- b. Clostridium botulinum
- c. Clostridium tetani
- d. Salmonella typhi
- 20. Which one is the example for qualitative analysis of milk?
- a. MBRT
- b. KOH mount
- c. LCB mount
- d. MPN test

# $PART - B (5 \times 5 = 25 \text{ Marks})$

Answer **ALL** questions

All questions carry equal marks

- 21. a) Write the contributions of Alexander Fleming (**OR**)
  - b) Write the contributions of Robert Koch.
- 22. a) Briefly explain bright field microscopy (**OR**)
  - b) Describe the principle and application of Dark field microscopy.
- 23. a) Explain the principle and steps involved in capsule staining (**OR**)
  - b) Explain the principle and steps involved in endospore staining.
- 24. a) Briefly explain the types of infection (**OR**)
  - b) Give a short note on Giardiasis.
- 25. a) Give a brief note on bioactive compounds (**OR**)
  - b) Write about the probiotics.

 $PART - C (3 \times 10 = 30 \text{ Marks})$ 

Answer ANY THREE questions

- 26. Describe the contributions of Louis Pasteur.
- 27. Describe the specimen preparation for Electron microscopy.
- 28. Explain Gram staining and acid fast staining.
- 29. Discuss in detail about Aspergillosis.
- 30. Explain in detail about the bacterial biopesticides.

# ALLIED PRACTICAL - I

Total Number of Hours: 30 3 Hours/ Week

## MICROBIOLOGY (PRACTICALS)

# **Course Objectives**

- To introduce the Microbiology laboratory
- To use the basic instruments in microbiology lab
- To study the morphology and movement of microbes
- To cultivate the microbes in laboratory
- To analyze the antibiotic susceptibility of microbes
- To detect the microbes from soil
- To ensure the quality of milk and water

#### **Course Outcome:**

CO1	The very basic laboratory practices and handling of hazardous material, biosafety importance, sterility and media preparations could be learned
CO2	These techniques would be very useful for quantitative analysis of microbes from environmental resources and also their physiological detection
CO3	Provides very essential procedure to separate / isolate pure culture from mixture of microorganisms and to study its physical characteristics
CO4	To get skilled in most common antibiotic sensitivity method and isolation of microbes from soil
CO5	Routine qualitative test for milk and water could be learned

- 1. Microbiology Good Laboratory Practices and Biosafety.
- 2. Preparation of culture media for bacterial cultivation.
- 3. Enumeration of bacteria from environment (soil/ water).
- 4. Staining techniques- simple, differential and negative staining.
- 5. Pure culture technique- Serial dilution, pour plate, spread plate and streak plate.
- 6. Determination of bacterial motility by hanging drop technique.
- 7. Antibiotic sensitivity test by Kirby Bauer method.
- 8. Isolation of microbes from rhizosphere soil.
- 9. Detection of quality of milk Resazurin, MBRT
- 10. Water Quality testing MPN.

# **Suggested Reading**

- 1. Cappucino J and Sherman N. (2010). **Microbiology: A Laboratory Manual**. 9<sup>th</sup> edition. Pearson Education Limited.
- P. Gunasekaran. (2005). Laboratory Manual in Microbiology. 1<sup>st</sup> Edition. New Age International Publishers.
- 3. Mette Praetorius Ibbe and Katherine Elasky. (2017). **Basic and Practical Microbiology Laboratory Manual**. 1<sup>st</sup> Edition. Cognella. Incorporated.
- 4. Norbel A.Tabo. (2004). Laboratory Manual in Microbiology. 1st Edition. Rex Book Store.
- 5. N.Kannan. (2002). **Laboratory Manual in General Microbiology**. 1<sup>st</sup> Edition. Panima Publishing Corporation.
- 6. Sundara Rajan. S. (2001). **Practical Manual of Microbiology**. 1<sup>st</sup> Edition. Anmol Publication Private

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2					
CO3		✓	✓	✓	
CO4	✓	✓			✓
CO5	✓	✓	✓	✓	✓

SEMESTER – III

18U3MBN01

Credits: 2

NMEC - I

Total number of Hours: 30

2 Hours/Week

## PUBLIC HEALTH AND HYGEINE

# **Course Objectives**

- i) To get an awareness about the public health and its significance
- ii) To gain the knowledge on the primary health care system in India
- iii) To provide an understanding of communicable and non communical diseases
- iv) To differentiate Occupational, Industrial and Urban Health

## **Course Outcome:**

CO1	The basic awareness on the public health and its significance could be learned
CO2	They could know the role of Primary health care system in India
CO3	The students were comprehend with the basic information about the communicable diseases
CO4	They could understand the basic information about the non-communicable diseases
CO5	They could aware of Occupational Safety & Health

UNIT - I Total No. of hours: 06

**Introduction to Public Health** – Introduction, Definition, Significance. Evolution of Public & community health. Determinants of Health – Biological, Behavioral, Socio-economic, Cultural, Environmental and Geographical.

UNIT - II Total No. of hours: 06

**Concept of Primary Health Care** – Public Health delivery system in India-Introduction to National Health Policy – 1983&2002, National Population Policy –2005, National Rural Health Mission (NRHM) and National Urban Health Mission (NUHM), National Public Health Programs.

UNIT – III Total No. of hours: 06

**Communicable & Infectious Diseases** – General overview of communicable diseases, impact of communicable diseases on developing.

UNIT - IV Total No. of hours: 06

**Non - Communicable Diseases -** Overview and introduction to NCDs-risk factors, prevention and management. General strategies, new approaches and policies of NCDs. NCDs programs of WHO, PAHO and Government of India.

UNIT-V Total No. of hours: 06

**Occupational, Industrial and Urban Health -** Occupational Safety & Health - Chemical and physical exposures, control of occupational exposures, injury control, occupational health disorders and diseases. Occupational health of working population of organized and unorganized sectors -Farmers, Industrial workers and health workers.

## **Suggested books**

- 1. Edward, Bouchieret and et al. (1995). Principles and Practice of Medicine. Davidson, Pearson Professional Ltd. London.
- 2. Jonathan Phillips, Paul Murray (1995). Biology of Disease. Black well Science Ltd. Australia.
- 3. Mackie and M.C. Cartney (1995). Practical Medical Microbiology. Longman Group, U.K.
- 4. David V. Mcqueen. Global Hand book On Non-Communicable Diseases and Health Promotion. Springer Publication.
- 5. S.L. Goel (2009). Education of Communicable and Non-Communicable Diseases. Deep & Deep Publications Pvt. Ltd.
- 6. David Vlahov, JoIvey Boufford, Clarence E. Pearson, Laurie Norris. Urban Health: Global Perspectives. Published by Jossey bass.
- 7. Jack E. Peterson (1991). Industrial Health American Conference of Governmental Industrial Hygienists.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2	✓		✓	✓	
CO3	✓	✓	✓	✓	✓
CO4	✓	✓			✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

# **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Third Semester Microbiology

	Wicrobiology
NMEC - P	UBLIC HEALTH AND HYGEINE
ime: Three Hours	Maximum Mark: 75
PA	ART - A (20  x  1 = 20  Marks)
	Answer <b>ALL</b> questions
All	l questions carry equal marks
1. Which of the following is a comr	nunicable disease?
a. Typhoid b. Dibeates	c. Obesity d. Heart diseases
2. Which of the following is a non of	communicable disease?
a. Malaria b. Measles	c. Typhoid d. Heart diseases
3. A longitudinal or prospective stu	udy is also referred to as an
a. ecological study b. cross-secti	
	ectious agent through the environment to a susceptible host
is called	
a. carrier b. reservoir	c. vector d. vehicle
5. Among the following, common o	occupational disease in working place
<u> </u>	n disease c. mental disease d. endocrine disease
6. The health services in any health	n care system may be of
a. Primary care level b. Secondary ca	
7. The function of any health care	
a. Production of resources	b. Management
c. Arrangement of resources into hea	<u> </u>
8. Health is best described as a reso	
a. A social and spiritual life	b. A productive social and economic life
c. Economic well-being	d. Physical capacity
9. What distinguishes primary hea	· · · · · · · · · · · · · · · · · · ·
a. A focus on primary, secondary and	- · · · · · · · · · · · · · · · · · · ·
b. Provision of interventions specific	·
c. Works within a multidisciplinary f	
d. Planning and operation of services	
	ygiene habit to teach young children?
a. Use a tissue to cover a sneeze	b. Don't share a glass or eating utensil
c. Wash hands frequently	d. Take a bath daily
* *	elated to change in health status defined as a
· ·	rminant c. Proximal determinant d. Ecological determinant
12. The main aim of public health	<u> </u>
a. Providing medical intervention app	propriate for the individual
b. Performing research to compare th	
c. Promoting health and preventing d	
d. Providing advice on risk markers a	* *
13. Ways to limit exposure to com	
	b. eating a balanced diet and participating in physical activity

d. All of the above

c. learn stress management techniques

14. Another word for communicable is ----a. harmless b. sudden c. contagious d. painful 15. Which of the following has not been associated with secondary brain injury? a. Hypoxia b. Hyperthermia c. Hyperglycemia d. Anemia 16. Choose which behavioral risk factors contribute to a person developing a NCD? a. Tobacco use b. Harmful use of alcohol c. Unhealthy diet d. Physical inactivity 17. The Occupational Safety and Health Act created three federal agencies for administration and enforcement. Which of the following is not one of them? a. The Occupational Safety and Health Administration (OSHA) b. The National Institute of Occupational Safety and Health (NIOSH) c. State Employee Health Commission (SEHC) d. The Occupational Safety and Health Review Commission (OSHRC) 18. The Secretary of Labor has authority to issue \_\_\_\_ involving new or improved techniques to safeguard safety or health of a worker. a. Temporary ordinances b. Experimental variances c. standard variances d. standing orders 19. The Occupational Safety and Health Act applies to all employees who work for an employer that is ----a. Internal security b. Interstate commerce c. Surrounding environment and natural ecosystems d. Overall health and safety of civilians. 20. Which body has the authority to order work to be halted? a. The government's Health and Safety Inspection Service b. The trade union c. The Arbodienst (Occupational Health and Safety Service) d. The supplier selection

## PART - B (5 x 5 = 25 Marks)

Answer **ALL** questions

All questions carry equal marks

- 21. a. Explain on evolution of public and community health (**OR**)
  - b. Write on biological and behavioral health determinants.
- 22. a. Give short note on natural and rural health machines (**OR**)
  - b. Describe national public health programmes.
- 23. a. Give an account on communicable diseases (**OR**)
  - b. What are all the impacts of communicable diseases?
- 24. a. Give an overview on non communicable diseases (**OR**)
  - b. Write a short note on occupational health disorders and diseases.
- 25. a. Give an introduction to NCDS risk factors. (OR)
  - b. Describe occupational safety

# $PART - C (3 \times 10 = 30 \text{ Marks})$

Answer **ANY THREE** questions

- 26. Explain in detail various health determinants.
- 27. Describe in detail on public health delivery system and its various policies.
- 28. Explain in detail overview on control of infectious and communicable diseases.
- 29. Give a detailed account on various risk factors and policies of non communicable diseases.
- 30. Describe in detail occupational health of working population in organized and unorganized sectors.



SEMESTER – IV

18U5MBC04

Credits – 4

Core – IV

Total number of Hours: 60

4 Hours/Week

#### IMMUNOLOGY AND IMMUNOTECHNOLOGY

# **Course Objectives:**

- 1. To gain knowledge about the cells and organs of the immune system.
- 2. To impart knowledge on immunity and vaccines.
- 3. To gain knowledge about antigens and immunoglobulins.
- 4. To impart knowledge on antigen-antibody interactions.
- 5. To understand about autoimmunity and hypersensitivity

## **Course Outcome:**

CO1	Structure and function of immune system and its importance in defense mechanism
	would be understood
CO2	It offers to understand immunological reactions / response and functions of immune
	cells
CO3	Ability to learn elaborative on antigen and antibody structure, reaction, activation and
	production of monoclonal antibodies
CO4	Helps to gain knowledge on antigen-antibody reaction and immunological tools for
	detection of causative agent
CO5	Concise immunological hypersensitivity and autoimmune disorders could be learned with background information

UNIT – I No. of Hours: 12

**Cells and organs of immune system:** Structure, function and properties of Lymphocytes, NK cell, Macrophage, Neutrophil, Eosinophil, Basophil, Mast cell, Dendritic cell - Primary lymphoid organs: Structure and function of Bursa, Bone Marrow and Thymus - Secondary lymphoid organs: Structure and function of Lymph Node, Spleen, GALT, MALT, CALT.

UNIT – II No. of Hours: 12

**Immune response:** Immunity - Concept of Innate and Adaptive immunity; Types - Specific and Non-specific - Primary and Secondary Immune Response; Generation of Humoral Immune Response (Plasma and Memory cells); Generation of Cell Mediated Immune Response (Self MHC restriction, T cell activation, Co- stimulatory signals) - Herd Immunity, Immunisation schedule, Vaccines - Definition and Types.

UNIT – III No. of Hours: 12

**Antigen, Antibody, MHC and Complement:** Antigen - Definition, types and characteristics - Haptens - Adjuvants. Immunoglobulins - Structure, Types, Functions and Properties - Theories of antibody synthesis - Hybridoma Technology and its Applications. Structure and Functions of MHC I & II molecules; Components of the Complement system; Activation pathways (Classical and Alternative pathways) - Biological consequences of complement Activation.

UNIT – IV No. of Hours: 12

Immunological Techniques: Principles and salient feature of Antigen-Antibody Interactions - Antibody affinity and avidity, Cross reactivity. Agglutination reactions - Blood grouping and Rh Typing, Haemagglutination, HAI, Bacterial agglutination, Passive agglutination. Precipitation reactions - in fluid and in gel. Immunoelectrophoresis, Immunofluorescence techniques - ELISA: Direct, Sandwich and Indirect, Biotin-Avidin system, RIA, Western blotting technique, Flowcytometry and Immunoelectron microscopy

UNIT – V No. of Hours: 12

**Immunological Disorders:** Hypersensitivity - Immediate and Delayed type Hypersensitivity - Gell and Coomb's classification of Hypersensitivity - Type I, II, III & IV - outline mechanisms with examples. Autoimmunity - Pernicious anaemia and Rheumatoid arthritis. Autoimune diseases.

## **Text Books**

- 1. Annadurai B (2008). A Textbook of Immunology and Immunotechnology. 1st Edition. S Chand & Co. Ltd., New Delhi.
- 2. Chakraborty P (2003).**A Text Book of Microbiology.** 2<sup>nd</sup> Edition. New Central Book Agency (P) Ltd, Kolkata.
- 3. Arti Kapil (2013). **Ananthanarayan and Paniker's Text Book of Microbiology**.9<sup>th</sup> Edition, Orient Blackswan Private Limited.

#### **Reference Books**

- 1. Kindt TJ, Goldsby RA, Osborne BA and Janis Kuby (2007). **Kuby Immunology.** W H Freeman and Company, New York.
- 2. Tizard IR (1995). **Immunology: An Introduction**. 4<sup>th</sup> Edition. Saunders College Publishers, USA.
- 3. Riott IM (1988). **Essentials of Immunology**, ELBS and Black Well Scientific Publishers, London

#### Web sources

- 1. https://nptel.ac.in/courses/102103038/1
- 2. https://nptel.ac.in/courses/102103038/39
- 3. https://nptel.ac.in/courses/102103038/download/module6.pdf
- 4. https://medlineplus.gov/ency/article/000821.html

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Fourth Semester

# IMMUNOLOGY AND IMMUNOTECHNOLOGY

Time:	Three Hours	Maximum 1	Marl	k: 7	5

# PART - A (20 x 1 = 20 Marks)

Answer **ALL** questions

	•	• •		
1. Macrophages are deri				
a) Monocytes b) L				
2. Cells which kills cells				
a) plasma cells b) re	ed blood cells	c) antigens	d) cytotoxic T lyr	nphocytes
3. Antibodies are secrete	d by			
a) Stem cells b) tiss	sue cells	c) plasma cells	d) membrano	us cells
4. Major components of	an immune syste	m include		
a) T-lymphocytes b)			d) All of Above	<u>,</u>
5. The two types of imm	unity in humans	are		
a) Intrinsic & Extrinsic		b) Innate & A d) Internal &	Acquired	
c) Overt & Covert		d) Internal &	External	
6. T cell mediates				
a) Cell mediated immune	e response	b) Humoral imr	nune response	
c) Non specific defence		d) None of thes	e	
7. Immunologic memory				
a) B cells b) T cell	ls c) Bot	h a & b	d) Phagocytes	
8. Origin & maturation of	of B cells takes pl	ace at		
a) Spleen b) Thyr	nus c) Bor	nemarrow	d) Lymphnodes	
9. A foreign macromoleo	cule that binds sel	lectively to an antil	body is called	
a) Stem cell b) An	tigen c) A	Antibody	d) Lymph	
10. IgM is structurally cl	haracterized as			
a) Monometric b)	Bimetric	c) Pentametric	d) Tetrametric	
11. The classical pathwa				
a) Bacteria, Antigen & A				
c) Antigen & Antibody	C	d) Virus, Antigen &	t Antibody	
12. MHC class I molecu				
a) Recognition of glycol	ipid antigens	b) Resistance to	fungi	
c) Resistance to viruses		d) Activation of	f neutrophils	
13. O blood group is uni				
a) Antigen A b) Ant	igen B c)	Antigen A & B	d) No antigens	
14. The precipitation test	t is relatively less	sensitive for the d	etection of	
a) Antigens b) Antibodie	es c) Complement	t d) Antigen-Antibo	ody complexes	
15. Commercially availa	ble ELISA kits a	re used for the dete	ection of	
a) Hepatitis B surface an				d) All of these
16. In 1959 radio immur	ie assay was deve	eloped by		
a) Soloman b) Be				

- 17. Inflammation reaction is brought about by -----
- a) Plasma cells
- b) Macrophages
- c) Mast cells
- d) Adipose cells
- 18. Which of the following binds to an Fc receptor on mast cells and basophils?
- a) IgA
- b) IgD
- c) IgM
- d) IgE
- 19. Pernicious anaemia develops from the deficiency of -----
- a) ATP
- b) Cobalt
- c) Hormones
- d) The intrinsic factors
- 20. What is the pathognomonic feature of rheumatoid arthiritis?
- a) Rheumatoid factor b) Rheumatoid nodule
- c) Morning stiffness
- d) ulnar drift of fingers

## PART - B (5 x 5 = 25 Marks)

Answer **ALL** questions

All questions carry equal marks

- 21. a) Describe about the Primary lymphoid organs (or)
  - b) Discuss about the History of immunology.
- 22. a) Write short notes on Specific immunity(or)
  - b) Explain about the T cell activation
- 23. a) Write short notes on Immunoglobulin structure (or)
  - b) Explain about MHC II molecules
- 24. a) Discuss about the Haemagglutination (or)
  - b) Give short notes on RIA (or)
- 25. a) Write short notes on Type I hypersensitivity reactions (or)
  - b) Discuss about the Type IV hypersensitivity reaction.

 $PART - C (3 \times 10 = 30 \text{ Marks})$ 

Answer **ANY THREE** questions

- 26. Briefly explain about the Haematopoiesis.
- 27. Explain about the generation of humoral immune response.
- 28. Discuss the Complement pathways.
- 29. Give a detailed account on ELISA.
- 30. Explain briefly about the autoimmunity.

CORE PRACTICAL – IV
Total number of Hours: 45
3 Hours/Week

## MAJOR PRACTICAL – IV - IMMUNOLOGY & IMMUNOTECHNOLOGY

## **Course Objectives:**

- 1. To know about the basics in immunology techniques
- 2. To get trained in the blood grouping
- 3. To gain knowledge in the agglutination tests
- 4. To understand the working principle and methods used in immunoelectrophoresis
- 5. To get skilled in diagnosis of various diseases through ELISA
- 6. To get trained in basics of complement fixation test

#### **Course Outcome:**

CO1	Able to perform ABO blood grouping and separation of serum and plasma
CO2	Can do latex agglutination tests and WIDAL
CO3	Ability to analyze antigen-antibody integration by immunoelectrophoresis
CO4	Trained with ELISA principle and procedure for the diagnosis of diseases
CO5	Can able to understand complement test

- 1. Identification of human ABO blood groups and Rh Typing.
- 2. Separation of serum/plasma from the blood sample (demonstration).
- 3. Latex agglutination test- RA Test, CRP Test, ASO Test.
- 4. WIDAL slide and tube agglutination technique.
- 5. Flocculation test RPR test.
- 6. Radial and ODD immunodiffusion technique.
- 7. Rocket immunoelectrophoresis.
- 8. Counter current immunoelectrophoresis.
- 9. Enzyme Linked Immunosorbent Assay (ELISA) demonstration.

## **References:**

1. Sambrook J and Russell DW (2001). **Molecular Cloning** - **A laboratory manual.** 3<sup>rd</sup> Edition. Cold Spring Laboratory Press, New York.

- 2. Surzycki S (2000). Basic Techniques in Molecular Biology. Springer-Verlag, New York.
- 3. Riott IM (1988). **Essentials of Immunology**, ELBS and Black Well Scientific Publishers, London.
- 4. Kindt TJ, Goldsby RA, Osborne BA and Janis Kuby (2007). **Kuby Immunology.** WH Freeman and Company, New York.
- 5. Chapel H and Halbey M (1986). Essentials of Clinical Immunology. ELBS, London.
- 6. Weir DM, Steward J (1993). **Immunology.** 7<sup>th</sup> Edition. ELBS, London.
- 7. Ausubel FM (1998). **Current Protocols in Molecular Biology.** Vol. 1 & 2. John Wiley & Sons Inc.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	<b>✓</b>	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓

SEMESTER – IV NMEC - II

18U4MBN02 Total number of Hours: 30 Credits: 2 2 Hours/Week

# **DISEASES - EPIDEMICS AND CONTROL**

## **Course Objectives:**

- i) To get an awareness about the infectious diseases and its epidemiology
- ii) To gain the knowledge on various chronic diseases
- iii) To impart the outbreak investigations and the role of laboratory
- iv) To endow with an geographic information system in infectious disease
- v) To aware the students in prevention of infectious diseases

#### **Course Outcome:**

CO1	They could learn about the infectious diseases and its epidemiological reports
CO2	They could learn the various chronic diseases
CO3	Able to learn about outbreak investigations and its diagnostic methods
CO4	Helps to gain the geographic informations in diseases
CO5	It makes expertise in the prevention of infectious diseases

UNIT I Total No. of hours: 06

**History of infectious disease and epidemiology** – Introductory concepts – Laboratory methods in the study of infectious diseases - Models to study infectious diseases - Modeling the spread of a disease - Emergent characteristics of Infectious diseases - Mathematical Epidemiology of Infectious disease.

UNIT II Total No. of hours: 06

**Chronic diseases** – common bacterial zoonotic diseases - Anthrax, Brucellosis – Vector borne diseases - Malaria and Dengue, Food borne Illness - Salmonellosis, Ameabiasis, Sexually Transmitted Diseases - HIV/ AIDS.

UNIT III Total No. of hours: 06

**Outbreak investigation** - Confirm outbreak and diagnosis - advance knowledge about a disease. Role of the Public Health Laboratory - Disease Surveillance - Principles of Screening and Screening Tests.

UNIT IV Total No. of hours: 06

**Geographic information systems in infectious disease** – Healthcare associated infections/infection prevention – Development of Drug Resistance & Infection Control in a Hospital Setting.

UNIT V Total No. of hours: 06

**Principles of elimination and eradication** – Vaccination. Behavior change and HIV/STDs - Blood Safety - Immigrant and Refugee Health - International Research in Resource Poor Settings - Critical Reading of Medical Literature

# **Suggested Reading**

- 1. Kenrad Nelson and Carolyn Williams (2014). Infectious Disease Epidemiology. Third Edition.
- 2. David L. Heymann (2015). Control of Communicable Diseases Manual. 20<sup>th</sup> Edition, American Public Health Association.
- 3. Annual Summary of Communicable Diseases Reported to the Minnesota Department of Health, 2015. Minnesota Department of Health.

## Web sources

- 1. www.health.state.mn.us/divs/idepc/newsletters/dcn/sum15/2015dcn.pdf
- 2. http://www.journals.uchicago.edu/CID/home.html

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓		✓	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5			✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

**----- / ----- 2018.** 

Fourth Semester Microbiology

**DISEASES – EPIDEMICS AND CONTROL Time: Three Hours Maximum Mark: 75** PART - A (20 x 1 = 20 Marks)Answer **ALL** questions All questions carry equal marks 1. In which of these do you see clue cells? a. Trichomonas vaginalis b. Bacterial vaginosis c. Candida d. HSV 2 2. In CSF of a patient with viral meningitis, the most prominent white cell is usually d. Polymorphs a. Monocytes b. Lymphocytes c. Eosinophils 3. Which is the most common organism/s causing osteomyelitis in all age groups? a. Streptococci b. Staphylococci c. Hemophilus d. Fungal 4. Which organism cannot be detected by antigen testing of CSF, serum of urine? a. Cryptococcus neoformans b. Mycobacterium tuberculosis c. E. coli d. Hemophilus 5. Which is not an AIDS defining illness? a. Oesophageal candidiasis b. PCP c. Pulmonary TB d. Invasive cervical cancer 6. Which is not a common cause of respiratory symptoms in HIV/AIDS patients? a. Community acquired bacterial pneumoniae b. Non Hodgkins lymphoma c. Pulmonary Embolus d. CMV 7. Which of these pulmonary conditions is most likely to be seen with a CD4 count between 200 and 500? a. Pulmonary TB b. CMV c. Karposi sarcoma d. Cryptococcus 8. Which is false regarding PCP pneumonia in AIDS? a. It is usually only seen when the CD4 count <200 b. Prophylaxis should be given in all pts with CD4 count <200 c. CXR characteristically shows bilateral diffuse infiltrates d. Once a patient has had it they are unlikely to get it again 9. Which statement is true regarding CT and LP in AIDS patients? a. They should all have a CT prior to LP b. If they are not febrile they do not need a CT b. If they have no focal neurology they do no need CT d. All of the above 10. Which vaccination should not be given to HIV suffers? a. ADT b. Pneumococcal c. DPT d. Inactivated polio vaccine 11. Which drug should not be given with midazolam? b. Lamivudine d. Ritinovir

c. Nevirapine

a. Zidovidine

12. Which drug regimen in AIDS is usually used?

c. 1 nucleoside, nevirapine and a protease inhibitor

a. 2 nucleosides and nevirapine

b. 2 nuclease and a protease inhibitor

d. All of the above

- 13. Which agent should not be part of the management of generalized tetanus?
- a. Metronidazole
- b. Penicillin
- c. Tetanus immunoglobulin
- d. Labetalol
- 14. A 60 year old lady presents with a skin tear to her left skin on her coffee table. She is unsure of her previous immunization status. How should this be managed?
- a. ADT
- b. immunoglobulin
- c. ADT and immunoglobulin
- c. None of the above

- 15. Which is not a differential diagnosis for tetanus?
- a. Strychnine poisoning
- b. Dystonic reactions
- c. Quinsy
- d. Cyanide poisoning

- 16. Which animal is least associated with rabies?
- a. Dogs b. Skunks
- c. Rats
- d. Bats
- 17. Which does not require post exposure prophylaxis for rabies?
- a. Scratch

- b. Bite on face
- c. Bite on extremity
- d. Skin contact with blood, urine or faeces
- 18. In which illness can hydrophobia be seen?
- a. Tetanus
- b. Malaria
- c. Rabies
- d. EBV
- 19. Which organism is least likely to show the characteristic periodicity of fever in malaria?
- a. P. malariae
- b. P. vivax
- c. P. ovale
- d. P. falciparum

- 20. Which statement is not true?
- a. Negative thick and thin smears does not adequately rule out malaria
- b. Falciparum malaria will always show up on thick and thin smears where the others may not
- c. Chloroquine is the drug of choice to treat falciparum
- d. Vivax and ovale are more likely to reactivate at a later stage

## PART - B (5 x 5 = 25 Marks)

Answer **ALL** questions

All questions carry equal marks

- 21. a) Explain in detail epidemiology of infection (OR)
  - b) Give short note on model study infections disease.
- 22. a) What are all the impacts of communicable diseases (OR)
  - b) Explain on pollutants reach humans.
- 23. a) Give short note on food borne illness (OR)
  - b) Give an overview on non communicable diseases.
- 24. a) Explain on evolution of public and community health (OR)
  - b) Give an account on communicable diseases.
- 25. a) Write on blood safety (OR)
  - b) Explain on occupational public health.

**PART** – **C** (3 X 10 = 30 Marks)

Answer **ANY THREE** questions

- 26. Explain and detail infectious disease and epidemiology.
- 27. Describe the concept of vector borne diseases.
- 28. Explain in detail about communicable diseases.
- 29. Give a detailed account on various risk factors and policies of non communicable diseases.
- 30. Give a detailed account on vaccination.

SEMESTER – IV **Total Number of Hours: 45 18U4MBA03** Credit - 4 4 Hours/ Week

## **MICROBIOLOGY**

# **Course Objectives:**

- To study the history of microbiology and to gain knowledge on microscopy
- To impart knowledge on bacterial anatomy and staining techniques
- To study the types of culture media, to understand sterilization techniques and to cultivate the microbes
- To understand the role of microbes in the field of medical, food and Environment

## **Course Outcome:**

CO1	Able to learn about chronological development and growth of microbiology and its
	importance and enables students to get motivated
CO2	It makes expertise in the art of techniques for the identification of microbes by
	staining methods
CO3	Enables to gather basic components of nutritional media, preparation and routine
	techniques used for the cultivation of microorganism in sterile condition
CO4	From this, one can stuff with medically and most prevalent diseases and its control /
	treatment
CO5	Helps to gain essential soil microbes and their significant role in agricultural field and
	food industry

UNIT – I No. of Hours: 09

History & Scope of Microbiology: Introduction - Contributions of various scientists to Microbiology - Louis Pasteur, Antony Van Leeuwenhoek, Robert Koch, Joseph Lister, Edward Jenner, Alexander Fleming. Microscopy: Bright field microscope, Dark field microscope, Phase contrast microscope, Fluorescent microscope and Electron microscope – TEM & SEM.

UNIT - II No. of Hours: 09

**Identification of Microbes**: Basic Structure of Bacteria – Gram positive and Gram negative bacteria. Stains and staining procedure - Types of staining - simple, differential and special staining – Fungal staining techniques – Lactophenol cotton blue staining and KOH mount.

UNIT - III No. of Hours: 09

**Cultivation of Microbes**: Culture media – Definition – Types - composition – Media preparation - Basal, Differential, Selective, Transport and Anaerobic culture media. Sterilization - Definition -Methods - Types of agents - Physical agents - Chemical agents. Culture techniques - Methods -Streak plate, Pour plate, Spread plate. Cultivation of anaerobes – Preservation of cultures.

ALLIED – III

UNIT – IV No. of Hours: 09

**Medical Microbiology:** Infection – Definition – Types – Mode of disease transmission – sources, Factors influencing pathogenesis – Disease cycle, Control of disease and prophylaxis. Peptic ulcer, Typhoid, Dengue, SARS, Candidiasis, Aspergillosis, Giardiasis.

UNIT – V No. of Hours: 09

**Applications of Microbiology:** Biofertilizer — Mycorrhiza, PGPR — Bioremediation — Biopesticides — Bacteria and Fungi, Biogas production - Bioactive compounds — Probiotics and prebiotics.

#### **Text Books**

- 4. Pelczar MJ, Chan ECS and Kreig NR (2008). **Microbiology**. 5<sup>th</sup> Edition, Tata McGraw Hill-Hill Education Pvt. Ltd., New Delhi.
- 5. Dubey RC and Maheswari DK (2005). **A Textbook of Microbiology**, Revised Multicolour Edition. S Chand and Company Limited, New Delhi.
- 6. Sullia S.B and Santhanam S (2005). **General Microbiology**. 2<sup>nd</sup> Edition, Oxford and IBH Publishing Co. Pvt. Ltd., New Delhi.

## **Reference Books**

- 4. Kathleen Park Talaro (2009). **Foundations in Microbiology: Basic Principles**, 7<sup>th</sup> Edition. McGraw-Hill Higher Education
- 5. Stanier RY, Ingraham JL, Wheelis ML and Painter PR (1987). **General Microbiology**. 5<sup>th</sup> Edition, MacMillan Education Ltd., London.
- 6. Gerard J Tortora, Berdell R Funke, Christine L Case (2010). **Microbiology: An Introduction.** 10<sup>th</sup> Edition, Pearson Benjamin-Cummings Publishing Company.

#### Web References

- 1. https://www.britannica.com/science/microbiology
- 2. https://www.atsu.edu/faculty/chamberlain/Website/Lects/Content1.htm
- 3. http://www.amm-mcrc.org/publications/Biofertilizers.pdf

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	<b>✓</b>
CO2			✓	✓	✓
CO3		✓	✓	✓	
CO4	✓	✓	✓		✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

# **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Second Semester Microbiology

ALLIED MICROBIOLOGY	
Time: Three Hours Maximum Mark: 75	
PART - A (20 x 1 = 20 Marks)	
Answer <b>ALL</b> questions	
All questions carry equal marks	
1. Robert Koch discovered	
a. Bacillus anthracis b. Salmonella typhi c. Ebola virus d. Amoeba parasite	
2. Anton Von Leeuwenhoek discovered	
a. Animalcules b. Virus c. Fungi d. Yeast	
3. Which one of the following is used to visualize live cells?	
a. Bright field microscopy  b. Dark field microscopy	
c. Phase contrast microscopy d. SEM	
4. Electron microscope was made by	
a. Robert hooke b. Knoll and Ruska c. Kepler and Galileo d. F.Janssen and Z.jansser	1
5. Gram staining is the example for	
a. Simple staining b. Differential staining c. Special staining d. None of the above	
6. Lipopolysaccharide is found in cell wall of	
a. Gram positive bacteria b. Gram negative bacteria c. Both d. Fungi	
7. Which one of the following is used as disinfectant in LCB staining?	
a. Lactic acid b. Phenol c. Glycerol d. Methylene blue	
8. Which of the staining technique helps in demonstrating spore structure in bacteria as well as fr	ee
spores?	
a. Acid-fast stain b. Endospore stain c. Capsule stain d. Flagella stain	
9. Which of the following is a rich source of nitrogen?	
a. Peptone b. Yeast extract c. Beef extract d. Agar	
10. The importance of agar in culture media were discovered by	
a. Ehrlich b. Petri c. Finly d. Hessy	
11. During preservation of microbes their	
a. characteristics change b. metabolism stop c. metabolism continue d. metabolism change	
12. Which of the following method is widely used for the preservation of microbes?	
a. Drying in vacuum b. Storage in sterile soil	
c. Lyophilization d. Storage in saline	
13. Transmission of 'pathogens' during pregnancy from mother to child is called as	
<ul><li>a. Direct transmission</li><li>b. Horizontal transmission</li><li>c. Vertical transmission</li><li>d. Indirect transmission</li></ul>	
14. An insect or animal carrier of disease is known as	

c. Fomite

d. Vehicle

b. Vector

a. Carrier

- 15. Water quality is measured by
- a. MBRT
- b. Resazurin test
- c. Staining
- d. MPN

- 16. Cholera is caused by
- a. Vibrio
- b. *E.coli*
- c. Salmonella
- d. Pseudomonas

- 17. Pasteurization technique is used for
- a. Milk
- b. Cheese
- c. Bread
- d. Antiseptic
- 18. The undesirable change that makes the unsafe food consumption is called -----
- a. Food decay
- b. Food spoilage
- c. Food loss
- d. All the above

- 19. Botulism is caused by
- a. E.coli
- b. Clostridium botulinum
- c. Clostridium tetani
- d. Salmonella typhi
- 20. Which one is the example for qualitative analysis of milk?
- a. MBRT
- b. KOH mount
- c. LCB mount
- d. MPN test

# **PART - B** (5 x 5 = 25 Marks)

Answer ALL questions

All questions carry equal marks

- 21. a) Write the contributions of Alexander Fleming (**OR**)
  - b) Write the contributions of Robert Koch.
- 22. a) Briefly explain bright field microscopy (**OR**)
  - b) Describe the principle and application of Dark field microscopy.
- 23. a) Explain the principle and steps involved in capsule staining (**OR**)
  - b) Explain the principle and steps involved in endospore staining.
- 24. a) Briefly explain the types of infection (**OR**)
  - b) Give a short note on Giardiasis.
- 25. a) Give a brief note on bioactive compounds (**OR**)
  - b) Write about the probiotics.

**PART – C** (3 X 10 = 30 Marks)

Answer **ANY THREE** questions

- 26. Describe the contributions of Louis Pasteur.
- 27. Describe the specimen preparation for Electron microscopy.
- 28. Explain Gram staining and acid fast staining.
- 29. Discuss in detail about Aspergillosis.
- 30. Explain in detail about the bacterial biopesticides.

## **ALLIED PRACTICAL - III**

Total Number of Hours: 30 3 Hours/ Week

# **ALLIED PRACTICAL - MICROBIOLOGY**

# **Course Objectives**

- To introduce the Microbiology laboratory
- To use the basic instruments in microbiology lab
- To study the morphology and movement of microbes
- To cultivate the microbes in laboratory
- To analyze the antibiotic susceptibility of microbes
- To detect the microbes from soil
- To ensure the quality of milk and water

## **Course Outcome:**

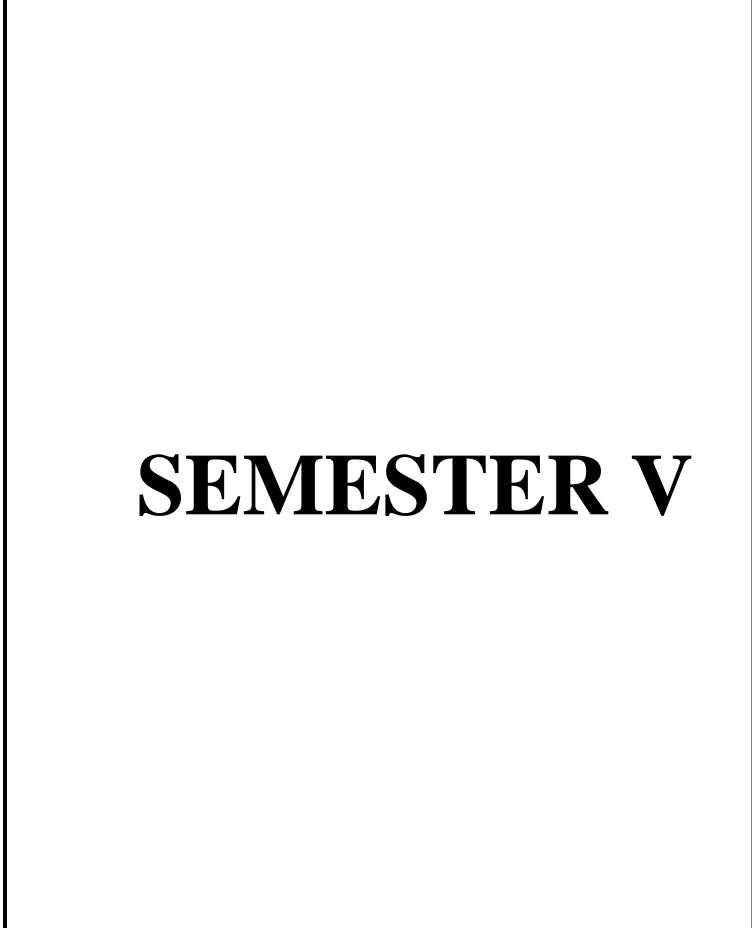
CO1	The very basic laboratory practices and handling of hazardous material, biosafety importance, sterility and media preparations could be learned
CO2	These techniques would be very useful for quantitative analysis of microbes from environmental resources and also their physiological detection
CO3	Provides very essential procedure to separate / isolate pure culture from mixture of microorganisms and to study its physical characteristics
CO4	To get skilled in most common antibiotic sensitivity method and isolation of microbes from soil
CO5	Routine qualitative test for milk and water could be learned

- 1. Microbiology Good Laboratory Practices and Biosafety.
- 2. Preparation of culture media for bacterial cultivation.
- 3. Enumeration of bacteria from environment (soil/ water).
- 4. Staining techniques- simple, differential and negative staining.
- 5. Pure culture technique- Serial dilution, pour plate, spread plate and streak plate.
- 6. Determination of bacterial motility by hanging drop technique.
- 7. Antibiotic sensitivity test by Kirby Bauer method.
- 8. Isolation of microbes from rhizosphere soil.
- 9. Detection of quality of milk Resazurin, MBRT.
- 10. Water Quality testing MPN.

# **Suggested Reading**

- 1. Cappucino J and Sherman N. (2010). **Microbiology: A Laboratory Manual**. 9<sup>th</sup> edition. Pearson Education Limited.
- 2. P. Gunasekaran. (2005). **Laboratory Manual in Microbiology**. 1<sup>st</sup> Edition. New Age International Publishers.
- 3. Mette Praetorius Ibbe and Katherine Elasky. (2017). **Basic and Practical Microbiology Laboratory Manual**. 1<sup>st</sup> Edition. Cognella. Incorporated.
- 4. Norbel A.Tabo. (2004). Laboratory Manual in Microbiology. 1st Edition. Rex Book Store.
- 5. N.Kannan. (2002). **Laboratory Manual in General Microbiology**. 1<sup>st</sup> Edition. Panima Publishing Corporation.
- 6. Sundara Rajan. S. (2001). **Practical Manual of Microbiology**. 1<sup>st</sup> Edition. Anmol Publication Private

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	<b>✓</b>	✓	✓
CO2		✓	✓	✓	✓
CO3		✓	✓	✓	
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓



SEMESTER – V CORE - V

18U5MBC05 Total number of Hours: 60 Credits: 6 6 Hours/Week

## MEDICAL BACTERIOLOGY AND MYCOLOGY

## **Course Objectives:**

- To study the pathogenesis, laboratory diagnosis and antimicrobial sensitivity testing
- To gain knowledge about the diseases caused by Gram positive and Gram negative cocci
- To impart knowledge on the diseases caused by Gram positive bacilli and Gram negative bacilli
- To understand the fungal classification, diagnosis, cultivation and antifungal agents
- To study the superficial, cutaneous, sub cutaneous, systemic and opportunistic mycoses

## **Course Outcome:**

CO1	Able to understand beneficial and harmful microbes
CO2	Medically important gram negative pathogens
CO3	Enterobacteria and other STI
CO4	Basics of fungal diseases and diagnostics methods
CO5	Dermatophytes and opportunistic mycosis

UNIT- I No. of Hours: 10

Normal microbial flora of human body – Infection – Types, Source, Modes of Transmission, Mechanism of bacterial pathogenesis – Collection and transport of clinical samples - Laboratory diagnosis of infectious diseases.

UNIT- II No. of Hours: 14

General characteristics, pathogenesis, clinical manifestation, laboratory diagnosis and control measures of the following pathogens - *Staphylococcus aureus*, *Streptococcus pneumoniae*, *Neisseria gonorrhoeae*, *Corynebacterium diphtheriae*, *Bacillus anthracis*, *Clostridium tetani* and *Mycobacterium tuberculosis*.

UNIT- III No. of Hours: 12

General characteristics, pathogenesis, clinical manifestation, laboratory diagnosis and control measures of the following pathogens - *Escherichia coli, Klebsiella pneumoniae, Proteus* species, *Salmonella typhi, Shigella dysenteriae, Pseudomonas aeruginosa, Vibrio cholerae, Treponema pallidum* and *Mycoplasma pneumoniae*.

UNIT- IV No. of Hours: 12

Classification of medically important fungi - Laboratory diagnosis of fungal diseases - Collection and examination of fungal specimens - Isolation and identification of fungi - Staining of fungi - KOH, LCB, PAS, H&E and GMS - Cultivation of fungi - Antifungal drugs mode of action - Antifungal susceptibility test .

UNIT- V No. of Hours: 12

Classification of Mycoses - Tinea nigra - Piedra - Dermatophytoses - Mycetoma - Histoplasmosis - Cryptococcosis - Candidiasis - Aspergillosis - Mycotoxicoses.

#### **Text Books**

- 1. Arti Kapil (2013). **Ananthanarayan & Jayaram Paniker's Text book of Microbiology**. 9<sup>th</sup> edition, Orient Longman Limited, Chennai.
- 2. Chakraborty P (2003). **A Text book of Microbiology.** 2<sup>nd</sup> edition, Published by New Central Book Agency (P) Ltd., Kolkata.
- 3. Jagdish Chander (2012). **Text book of Medical Mycology**. 3<sup>rd</sup> edition. Mehta Publishers, New Delhi.

#### **Reference Books**

- 1. Jawetz E and JL Melnic (2001). **Medical Microbiology**, 22<sup>nd</sup> edition, Tata Mc Graw-Hill, New Delhi.
- 2. David Greenwood CB and Richard (2002). **Medical Microbiology**. 22<sup>nd</sup> edition, Tata Mc Graw- Hill, New Delhi.
- 3. Monica Cheesbrough (2003). **District Laboratory Practice in Tropical Countries**. Part 1 and 2. Low-Price edition, Cambrige University Press.

#### Web sources

 $https://www.cartercenter.org/resources/pdfs/health/ephti/library/lecture\_notes/med\_lab\_tech\_students/ln\_med\_bact\_final.pdf$ 

https://mycology.adelaide.edu.au/mycoses/

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017 - 18 onwards)

# **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Fifth Semester

Microbiology

## MEDICAL BACTERIOLOGY AND MYCOLOGY

Maximum Marks: 75
d. Streptococcus
a. s ep rece cens
and in
d. All of above
ser's stain
d. None of the above
d. Acid fast bacilli
stridium difficile
bit
n
e bile salt agar
ophilus ducryi
streptobacillus
01-1
Rickettsiae
d. proteins
ts in single form
is in single lottii
d. Von Kossa
G. VOII IXOSSU

a. Allylamines c. Echinocandis d. Thiocarbamate b. Polynes 17. Which of the following is not the characteristic of histoplasmosis? a. Person to person transmission b. Specific geographic distribution c. Yeasts in tissue d. mycelial phase in the soil 18. Causative agent for 'ringworm' is ----a. Epidermatophyton b. *Tinea nigra* c. Mycetoma d. Histoplasma 19. Tinea pedis' is scientific name of a foot disease that is commonly called as a. Athlete's foot b. Ringworm c. Skin rash d. Skin infection 20. Which one is considered as class one carcinogen?

> **PART - B**  $(5 \times 5 = 25 \text{ Marks})$ Answer **ALL** the Questions All questions carry equal marks

c. Fumonisin

d. Ochratoxin

21. a) What are the types of infections? (OR)

a. Aflatoxin

- b) Briefly explain the antibacterial susceptibility testing.
- 22. a) What are the virulence factors involved in Staphylococcal infections? (OR)

b. Ergotoxin

- b) Discuss the different types of anthrax.
- 23. a) Write a note on different kinds of diarrhea caused by *E.coli*. (OR)
  - b) Describe about cholera and its diagnosis.
- 24. a) Explain about classification of medically important fungi (OR)
  - b) Write a short note on systemic mycosis samples
- 25. a) Explain about *Tinea nigra* (OR)
  - b) Write a short note on Histoplasmosis

**PART - C**  $(3 \times 10 = 30 \text{ Marks})$ Answer **ANY THREE** the Questions All questions carry equal marks

- 26. Explain in detail the normal flora of the human's different anatomical sites.
- 27. Write a detailed note on morphology, virulence factors and pathogenicity of *Clostridium perfringens*.
- 28. Write the pathogenesis and laboratory diagnosis of Syphilis.
- 29. Explain about Antifungal susceptibility test.
- 30. Write a short note on Candidiasis and cryptococcosis.

SEMESTER – V

18U5MBC06

Credits: 5

CORE - VI

Total number of Hours: 60

5 Hours/Week

### INDUSTRIAL AND PHARMACEUTICAL MICROBIOLOGY

## **Course Objectives:**

- To gain knowledge about screening techniques and strain improvement.
- To study about different types of bioreactors.
- To know about industrial production of enzymes and antibiotics.
- To understand the types of pharmaceutical products.
- To study the quality control of pharmaceutical products.

## **Course Outcome:**

The students could able to gain knowledge on

CO1	Basic background information on industrial strain development and sterilization
CO2	Downstream and upstream process of production technology
CO3	Various industrial microbiological product synthesis
CO4	Synthesis of antimicrobial drugs using fermentation technology
CO5	Drug delivery mechanism and clinical trials

UNIT - I No. of Hours: 12

**Introduction to industrial microbiology:** Industrially important microorganisms - Screening techniques - Primary and Secondary. Upstream processing - Strain improvement - Development of inoculums – Production media – Industrial sterilization.

UNIT - II No. of Hours: 12

Industrial Fermentor - Components of fermentor - Types of bioreactors - Types of fermentor instrumentation –Scale up of fermentation - Down Stream Processing – Recovery and Purification of intracellular and extracellular products and natural sources.

UNIT - III No. of Hours: 12

Industrial production of enzymes  $-\alpha$  amylase. Organic acid - citric acid, lactic acid and acetic acid. Alcoholic beverages - Wine and Beer. Aminoacid - glutamic acid. Vitamin - B12. Microbiological production of antibiotics - Penicillin and streptomycin.

UNIT - IV No. of Hours: 12

Types of pharmaceutical products - Antimicrobial agents - Bioassay of antimicrobial agents - Contamination, spoilage and preservation of pharmaceutical products - Microbiological quality control - Sterility test- Pyrogen test- Toxicity test- Carcinogenicity test.

UNIT - V No. of Hours: 12

Drug delivery systems - Drug distribution in body - Bio-availability- Adverse drug reaction and drug interaction. Drug discovery - Phases of drug discovery - Clinical studies: phase I and phase II of clinical trials - Bioprospecting - Extraction, purification and characterization of bioactive molecules from natural resources.

#### **Text Books**

- 1. Patel A.H (2011). **Industrial Microbiology**. 2<sup>nd</sup> edition. Published by Mac Millan Publishers India Ltd., Chennai.
- 2. Cassida L.E (1996). **Industrial Microbiology**. New Age International Publishers, Chennai.
- 3. Purohit S.S, Saluja A.K and Kakrani H.N (2004), **Pharmaceutical Microbiology**, 1<sup>st</sup> edition, Agrobios (India), Jodhpur.

## Reference books

- 1. Peppler H.J and Perlman D (1979). **Microbial Technology**. Vol.1 and II. 2<sup>nd</sup> edition. Academic Press, New York.
- 2. Stanbury P.F, Whitaker A and Hall S.J (1995). **Principles of Fermentation Technology**. 2<sup>nd</sup> edition. Pergamon Press, New York.

#### Web Sources

https://pdfs.semanticscholar.org/635d/da50cbf522a7c860ddf899925ffa703123b1.pdf https://run.edu.ng/directory/oermedia/422231995398.pdf

http://site.iugaza.edu.ps/mwhindi/files/Modern-Industrial-MicrobiologyBiotechnology.pdf

file:///H:/industrial/0c03ce4cbbae680f46362dd24207e254-original.pdf

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓			✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	
CO4	✓	✓	✓	✓	
CO5	✓		✓	✓	✓

18U5MBC06

(For the candidates admitted from 2017 - 18 onwards)

# **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Fifth Semester

Microbiology

## INDUSTRIAL AND PHARMACEUTICAL MICROBIOLOGY

<b>Time: Three hours</b>		Maximum Marks: 75
	<b>PART - A</b> $(20 \times 1 = 20 \text{ Marks})$	
	Answer <b>ALL</b> the Questions	
	All questions carry equal marks	

1. The best medium for the production of Penicillin is
a. Nutrient agar b. Corn steep liquor c. Sulfite waste liquor d. Whey
2. Industrially important Antibiotic producing organisms shall be isolated by
a. Disk plate method b. Direct plate method c. Serial dilution method d. Crowded plate method
3. In the industrial production of streptomycin, the secondary metabolite or by products is
a. Vitamin – B12 b. Vitamin – C c. Vitamin – B6 d. Ethanol
4. The fungus used in the industrial production of citric acid:
a. Rhizopus Oryzae b. Fusarium moniliformae c. Rhizopus nigricans d. Aspergillus nigricans
5. Vitamin B12 can be estimated and determined by using organism
a. Lactobacillus spp b. Lactobacillus Leichmanni c. Bacillus subtilis d. E. coli
6. Batch fermentation is also called
a. Closed system b. Open system c. Fed-Batch system d. Sub-merger system
7. Industrial microbiology, mainly depends on the phenomenon
a. Pasteurisation b. Fermentation c. Vaccination d. Purification
8. For thorough mixing of medium and inoculum the part of fermentor useful is
a. Shaft b. Headspace c. Impeller d. Sparger
9. Different methods of strain improvement are
a. Protoplast fusion b. Recombinant DNA technique c. Genetic recombination d. All of these
10. The purification and recovery of the production after fermentation is called
a. Upstream process b. Downstream process c. Surface fermentation d. None of these
11. Who developed the concept of specific toxicity?
a. Pasteur b. Fleming c. Watson d. Ehrlich
12. The susceptibility of a microorganism to antibiotics and other chemotherapeutic agents can be
determined by using
a. tube dilution technique b. paper disk plate c. both (a) and (b) d. none of these
13. The process that is used to prove that a drug is safe and effective in treating specific conditions in
certain patient populations?
a. Drug discovery b. Preclinical development c. The patent process d. Clinical development
14. Which of the following packaging material is protect the drug content against light
a. Plastic containers  b. Amber colored glass containers
c. metal containers d. all of the above

- 15. What is the purpose of pre-clinical testing?
- a. To verify that a drug is sufficiently safe and effective to be tested in humans.
- b. To undergo preliminary testing in healthy humans to monitor the effects of the drug.
- c. Both a and b
- d. To create a basic outline for the larger scale future tests on a widespread population.
- 16. On what do Phase 2 clinical trials test?
- a. Animals

- b. Large-scale tests in people with the target disease/population
- c. People with the target disease/condition
- d. Healthy human volunteers
- 17. Bioprospecting is -----
- a. the search for gold from marine sources
- b. the search for fuel in sea
- c. the search for pharmacological or other chemicals from natural resources
- d. none of the above
- 18. In fermenter, up to the production of desirable product is termed
- a. Upstream process
- b. Downstream process
- c. Fermentation
- d. Sterilisation
- 19. Which method of purification allows separation of solids from fluids (liquids or gases) by interfering a medium through which only the fluid can pass?
- a. Filtration b. Precipitation
- c. Centrifugation
- d. Sedimentation
- 20. Which separation technique is based on differential partitioning between two phases that is mobile and stationary?
- a. Filtration
- b. Precipitation
- c. Centrifugation
- d. Chromatography

## **PART - B** $(5 \times 5 = 25 \text{ Marks})$

Answer **ALL** the Questions

All questions carry equal marks

- 21. a). Write a short note on auxanography technique (OR)
  - b). List out the different methods of inoculumdevelopment.
- 22. a). Give an account on upstream process (OR)
  - b). Write in detail about different types of fermentor.
- 23. a). Elaborate the protocol for the industrial production of  $\alpha$ -amylase (OR)
  - b). Write a note on the production of citric acid
- 24. a). Explain the bioassay of antimicrobial agents? (OR)
  - b). Write about the contamination, spoilage and preservation of pharmaceutical products? (OR)
- 25. a). Write short notes on phases of drug discovery
  - b). Explain the drug distribution in body?

## **PART - C** $(3 \times 10 = 30 \text{ Marks})$

## Answer **ANY THREE** the Questions

All questions carry equal marks

- 26. Give an account on the various methods of screening the industrially important Microorganisms
- 27. Draw a neat sketch of a fermentor and explain in detail about its parts.
- 28. Write an account on the industrial production of acetic acids
- 29. Describe the phases of clinical studies
- 30. Explain about the drug distribution in body?

SEMESTER – V

18U5MBC07

Credits: 5

CORE - VII

Total number of Hours: 60

5 Hours/Week

#### **GENETIC ENGINEERING**

### **Course Objectives:**

- 1. To get hold of knowledge on enzymes and vectors
- 2. To be familiar with rDNA technology
- 3. To obtain knowledge about molecular techniques
- 4. To know the basics on genetic engineering in plants
- 5. To obtain knowledge in the basics on genetic engineering in plants

## **Course Outcome:**

The students could expertise in

CO1	Restriction modification system and vectors
CO2	Natural gene transfer methods
CO3	Molecular genome amplification techniques
CO4	Use of bacterial Ti, Ri plasmids and plant gene targeting techniques
CO5	Transgenic technology and animals

## **UNIT - I Restriction Enzyme and Vectors**

Restriction and modification System in Bacteria (*E.coli*) - Restriction endonucleases type I, II & III. Vectors - Plasmids - Phage, Cosmids, Phagemids, special vectors-broad host range expression in bacteria, shuttle vectors.

#### **UNIT - II Gene Recombination and Gene transfer methods**

Bacterial conjugation – transformation – transduction, Microinjection, Electroporation, gene Gun method, Ultrasonication, Liposome fusion, Microlaser gene transfer

## **UNIT - III PCR and Its applications**

PCR technology-gene amplification- PCR primer designing and optimization; variation in PCR (RT PCR, RACE), RAPD, RFLP.

No. of Hours: 12

No. of Hours: 12

No. of Hours: 12

### **UNIT - IV Genetic engineering in plants**

Uses of *Agrobacterium tumefaciens* and *Arhizogenes*, Ti plasmids, Strategies for gene transfer to plant cells, Direct DNA transfer to plants, Gene targeting in plants, Use of plant viruses as episomal expression vectors.

## **UNIT - V Genetic engineering in animals**

Production and applications of transgenic mice, role of ES cells in gene targeting in mice, Therapeutic products produced by genetic engineering-blood proteins, human hormones, immune modulators and vaccines.

## **Suggested Reading**

- 1. Clark DP and Pasternik NJ. (2009). Biotechnology: Applying the Genetic Revolution. Elsevier Academic Press, USA.
- 2. Brown T.A (2010). Gene cloning and DNA Analysis. 6<sup>th</sup> edition. Blackwell publishing, Oxford, U.K.
- 3. Satyanarayana U 2005 Biotechnology 1<sup>st</sup> edition. Books & Allied (p) Ltd.-Kolkata.
- 4. Primrose SB and Twyman RM. (2006). Principles of Gene manipulation and Genomics. 7<sup>th</sup> edition, Blackwell publishing, Oxford, U.K.
- 5. Dubey R. C. A Textbook of Biotechnology. Publisher: S. CHAND.
- 6. Primonrose SB and Twyman RM. (2008). Genomics: Application in human biology Blackwell publishing, Oxford, U.K.

#### **Web Sources**

https://nptel.ac.in/downloads/102103013/

https://science.umd.edu/classroom/bsci124/lec41.html

http://genok.no/wp-content/uploads/2013/04/Chapter-4.pdf

#### **Mapping**

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	<b>✓</b>	✓	✓	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5		✓	✓		✓

No. of Hours: 12

No. of Hours: 12

(For the candidates admitted from 2017 - 18 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Fifth Semester

Microbiology

#### **GENETIC ENGINEERING**

Time: Three hours **Maximum Marks: 75** 

> **PART - A**  $(20 \times 1 = 20 \text{ Marks})$ Answer ALL the Questions All questions carry equal marks

- 1) Thymosin proved effective against brain an
- a) Liver cancer
- b) Stomach cancer
- c) Lung cancer
- d) Blood cancer

- 2. What is the final product of the RNase H method?
- a) blunt ended dsDNA
- b) staggered dsDNA at both ends

d) DNase

- c) staggered dsDNA at 3' end
- d) staggered dsDNA at 5' end
- 3. What would not happen if the RNA strand is completely removed from RNA: DNA hybrid?
- a) There are no chances of the synthesis of the second DNA strand
- b) Chance complementarity would take place
- c) Hairpin structure would be formed
- d) Hairpin structure is formed is not the final structure
- 4. The loop region is single stranded. It can be cleaved by using which enzyme?
- 5. Choose the correct statement with respect to the self priming method of cDNA synthesis.
- a) It is less preferred than RNaseH method
- b) A hairpin structure is formed with guarantee

a) Exonuclease b) S1 nuclease c) RNaseH

- c) The sequence corresponding to the 5' end is lost
- d) Reverse transcriptase is not used
- 6. Choose the incorrect statement for the method homopolymer tailing.
- a) The first step is the RNA: DNA hybrid synthesis
- b) Terminal transferase is used for the addition of nucleotides on 3' end
- c) Terminal transferase adds only at DNA strands
- d) The DNA strand is now having known sequence at 3' end
- 7. Choose the correct statement for RACE.
- a) It stands for Random Amplification of cDNA ends
- b) It is for cloning particular cDNA ends
- c) It is only of one type, which is 5' RACE
- d) Sequence data is not available in any case
- 8. The first primer in the case of 3' RACE is
- a) internal sequence
- b) oligo-dT adaptor molecule
- c) oligo-dA adaptor molecule
- d) adaptor oligo-dT primer
- 9. The first cDNA strand in 5' RACE is tailed with oligo-dA tail.
- a) True
- b) False

10. What is the second primer in the case of 5' RACE?  a) Internal primer b) Oligo-dA sequence c) Adaptor-oligo-dT primer d) Oligo-dT adaptor molecule 11. A vector is a plasmid used to transfer the a) Chromosome b) Gene c) Nucleus d) Cell
12) gene therapy healthy gene are used to replace a) Dead gene b) Abnormal gene c) Defective gene d) Old gene
13) Genetic engineering increases effeciency and
a) Productivity b) Metabolism c) Meiosis d) Mitosis
14) A genetic code is a message store in
a) Cell b) Nucleus c) Cytoplasm d) Gene
15) Fermenter are used to culture
a)Algae b) Fungi c) Virus d) Bacteria
16) Process of manipulating genes usually outside normal reproductive process is known as
a) genetic modification b) gene targeting c) genome recombination d) gene
linking 17) First genetically modified organism generated was
a)Fish b) bacteria c) mice d) virus
18) First genetically modified mice is generated in
a) 1968 b) 1964 c) 1974 d) 1978
19) First genetically modified pet was sold in United States in
a) 2003 b) 2008 c) 2006 d) 2004
20) Enzyme which is used to remove or knockout genes is known as
a) Nucleolus b) nuclease c) nucleotide d) clones
<b>PART - B</b> $(5 \times 5 = 25 \text{ Marks})$
Answer <b>ALL</b> the Questions
All questions carry equal marks
21. a) Describe enzymes used in genetic engineering (or)
b) Discuss bacterial conjugation.
22. a) Write short notes on microinjection (or)
b) Explain ultrasonication.
23. a) Write short notes on isolation of plasmid DNA (or)
b) State about transformation and transfection.
<ul><li>24. a) Write the significant application of PCR (or)</li><li>b) Give short notes on Ti plasmids.</li></ul>
25. a) Write short notes on gene technology in medicine (or)
b) Discuss transgenic animal and its application.
3/ = 32 20 20 20 20 20 20 20 20 20 20 20 20 20
<b>PART - C</b> $(3 \times 10 = 30 \text{ Marks})$
Answer ANY THREE Questions
All questions carry equal marks
26. Write an essay on production and application of transgenic mice?
27. Explain restriction endonucleases type I and type II.
28. Elaborate the various methods of Microlaser gene transfer technology.

29. Give a brief account on PCR and its application.

30. Explain briefly about transformation

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SEMESTER – V 18U5MBCP05 Credits: 3 CORE PRACTICAL - V
Total number of Hours: 45
6 Hours/Week

#### PRACTICAL V

## **Course Objectives:**

- To obtain knowledge about fungal identification methods
- To gain information about immobilization technique
- To know the techniques in amylase production from bacteria
- To update the identification methods used in clinical pathogen detection
- To get knowledge about citric acid producing fungi

#### **Course Outcome:**

They students could able to do

CO1	Diagnosis of pathogens from clinical samples
CO2	Demonstration of fungal pathogens
CO3	Screening of bacteria for amylase production
CO4	Screening of bacteria producing citric acid
CO5	Immobilization of products for preservation

- 1. Identification and Biochemical characterizations of clinical pathogens from clinical samples Urine, Pus and Sputum.
- 2. Identification of fungal specimens by direct microscopy KOH and LCB preparations.
- 3. Screening of amylase producing bacteria from soil.
- 4. Screening of citric acid producing bacteria from soil sample.
- 5. Immobilization technique.
- 6. Isolation of plasmid DNA from E. coli.
- 7. Screening of recombinants Blue / white selection assay.

#### **Suggested Manuals**

- 1. Arora, B and D.R. Arora, (2013), **Practical Microbiology** CBS Publishers & distributors Pvt. Ltd, New Delhi.
- 2. Benson, J.H., (2001), "Microbiological Applications: A Laboratory Manual in General Microbiology", Eighth Edition, McGraw-Hill, New York.
- 3. Cappuccino, J.G. and N. Sherman, (2005), "Microbiology A Laboratory Manual", Seventh Edition, Benjamin and Cummings Publications, San Francisco.

- 4. Gunasekaran, P., (2005), "**Laboratory Manual in Microbiology**", New Age International (P) Ltd, New Delhi.
- 5. Kannan, N., (2003), "**Laboratory Manual in General Microbiology**", Fourth Edition, Palani Paramount Publications, Palani.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1					
CO2					
CO3					
CO4					
CO5					✓

SEMESTER – V

17U5MBE01

Credits: 4

ELECTIVE - I

Total number of Hours: 45

4 Hours/Week

#### HAEMATOLOGY AND BLOOD BANKING

## **Course Objectives**

- To gain knowledge about the blood cells.
- To study hematological diseases.
- To impart knowledge on hematological tests.
- To gain knowledge about immunohematology.
- To study blood banking and blood transfusion.

#### **Course Outcome:**

CO1	Basics of hematology and immune cells
CO2	Immunological and deficiency-oriented disorders
CO3	Analysis of cells by various mathods
CO4	Routine hematological tests
CO5	Blood transfusion and disease transfer

UNIT - I No. of Hours: 09

Introduction to Haematology – Blood – Components and its function - Haematopoietic system of the body – Development of Blood corpuscles - Erythropoiesis – Leukopoiesis – Thrombopoiesis

UNIT - II No. of Hours: 09

Haematological diseases – Anaemia - Types of Anaemias – Iron deficiency anemia. Haemolytic disease of the new born – Infectious Mononucleosis – Multiple myeloma – Parasitic infections of blood – Leukaemia - classification.

UNIT - III No. of Hours: 09

Routine haematological tests – Introduction – Collection of blood – Anticoagulants - Complete Blood Cell count (CBC) – Determination of Haemoglobin by Sahli's method – Cynamethaemoglobin method – RBC count – WBC count - Differential count – Determination of ESR.

UNIT - IV No. of Hours: 09

Haemostasis and Blood Coagulation – Mechanism of coagulation – Determination of Bleeding time and Clotting time – Immunohaematology – Human blood group systems – ABO grouping and other blood group systems – Rh Typing.

UNIT - V No. of Hours: 09

Blood banking and Blood transfusion – Screening of blood donors – Preservation and storage of donated blood - Cross matching – Blood transfusion – HLA typing - Transfusion transmitted diseases – Transfusion reaction.

#### **Text Books**

- 1. Drew Provan (2009). ABC of Clinical Haematology, 3<sup>rd</sup> edition. BMJ books.
- 2. Hoffbrand A.V, Pettit J.E and Moss P.A.H (2001). Essential Haematology. 2<sup>nd</sup> edition. Blackwell Science, New York.

#### **Reference Books**

- Denise M Harmening (2012). Modern Blood Banking and Transfusion Practices. 6<sup>th</sup> Edition. F A Davis Company, Philadelphia.
- 2. Transfusion Medicine Technical Manual (2003). 2<sup>nd</sup> edition. DGHS, Ministry of Health and Family Welfare, Govt. of India,
- 3. Peter Delves, Seamus Martin, Dennis Burton (2006). Roitt's Essential Immunology. 11<sup>th</sup> edition. Wiley-Blackwell, New York.

#### Web sources

https://nptel.ac.in/courses/102103012/pdf/mod7.pdf

 $https://www.cartercenter.org/resources/pdfs/health/ephti/library/lecture\_notes/med\_lab\_tech\_students/ln\_hematology\_mlt\_final.pdf$ 

http://www.rajswasthya.nic.in/RHSDP%20 Training%20 Modules/Lab.%20 Tech/Blood%20 Banking/Introduction.pdf

file:///H:/Hematology/abo%20blood%20grouping.pdf

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Fifth Semester Microbiology

## HAEMATOLOGY AND BLOOD BANKING

	IIAEMATOLOGI	AND DECOD DAN	IXII VO	
<b>Time: Three Hours</b>			Maxim	num Mark: 75
	PART - A	(20  x  1= 20  Marks)		
	Answer	<b>ALL</b> questions		
	All question	is carry equal marks		
1. The process of platelet	•	• •		
	B. Thrombopoiesis		D Fryth	ropoiesis
2. Component of the red b	-	-	•	iropolesis
	B. Nucleus	C. Plasma	D. Blast	cell
3. Blood test that indicate				
A. Bone marrow biops:				oifferential count
4. Hormone that stimulate				
	B. Interleukin	C. Erythropo	oietin I	O. Insulin
5. Lymphopoiesis occur	in	, ,		
A. Lymphatic tissue	B. Lymphocyte	es C. Bone mar	row I	O. Plasma
6. Is concerned primarily	with phagocytosis			
A. Antibodies	B. RBC	C. Platelets	D. WBC	
7. Is decreased in anemia				
A. Platelets	B. Stem cells	C. RBC	D. WBC	
8. Primarily concerned w			<b>.</b>	
A. Platelets	B. WBC	C. Erythrocyte	D. Haemogl	obin
9. Involved in a hemolytic			N D1 4 4	
A. WBC	B. RBC		D. Blastocyte	
10. Enzyme that converts	-	moin C. Prothrombin activ	votom D II	listamina
A. Heparin 11. An anticoagulant that				listamine
A. Coumadin		<u> </u>	D. Thrombus	
12. Enzyme that activates	1		J. TillOllious	
A. Thrombus		C. Platelets	D. Thrombia	n
13. The universal recipie		C. I lacolocis	D. Timomon	•
A. A+		C. B+ D. AE	3+	
14. The plasma of this bl	ood type contains bo	oth anti-A antibodies	and anti-B ar	ntibodies
	B. O- (			
15. A person with this blo	od type can receive	(by transfusion) only	y type O- bloo	od
A. AB+ B.	O- C. B+	D. O+		
16. The positive and nega		· ·	gen	
	•	n factor D. AE	BO grouping	
17. Reticulocytes are usua	2			
A. Sickle cell anemia	B. Aplastic and	emia C. Iron defici	ency anemia	D. Leukemia

- 18. A megaloblastic anemia that is treated with vitamin B12 injections
  - A. Pernicious anemia

B. Sickle cell anemia

C. Aplastic anemia

- D. Iron deficiency anemia
- 19. An individual who has recently been diagnosed with syphilis is deferred for:
  - A. 4 weeks
- B. 2 weeks

C. Permanently

D. 1 year

20. Red blood cells can be frozen and stored up to:

A. 3 years

B. 5 years

C. 7 years D. 8 years

PART - B (5 x 5 = 25 Marks)

Answer **ALL** questions

All questions carry equal marks

- 21. (a). Write notes on Blood components (**OR**)
  - (b). Give the details about the Erythrpoiesis
- 22. (a). Discuss about Infectious Mononucleosis (OR)
  - (b). Explain about Blood parasitic infections.
- 23. (a). Give an account on Anticoagulants (**OR**)
  - (b). Discuss on Differential count
- 24. (a). Give an account on Determination of Clotting time (**OR**)
  - (b). Explain about ABO grouping.
- 25. (a). Account on Preservation and storage of donated blood. (OR)
  - (b). Explain the HLA typing.

**PART – C** (3 X 10 = 30 Marks)

Answer **ANY THREE** questions

All questions carry equal marks

- 26. Haematopoietic system of the body? Explain.
- 27. Explain about Leukaemia and its classification.
- 28. Determination of Haemoglobin by Sahli's method Cynamethaemoglobin method.
- 29. Explain in detail on Mechanism of Blood coagulation.
- 30. Explain the Transfusion reactions.

SEMESTER – V

18U5MBE02

Credits: 4

ELECTIVE - I

Total number of Hours: 45

4 Hours/Week

### ENTREPRENEURSHIP IN MICROBIOLOGY

## **Course Objectives**

- To understand the basic concepts of entrepreneurship and become a young women entrepreneur.
- To gain business opportunities on mushroom cultivation.
- To expand systemic knowledge on different composting technology.
- To increase the comprehension on various biotechnological approaches to establish successful enterprises.
- To understand different financial agencies supporting entrepreneurship.

#### **Course Outcome:**

CO1	Entrepreneur importance towards women development
CO2	Mushroom cultivation and various products development
CO3	Bio-composting and its application
CO4	Biofertilizer manufacturing techniques
CO5	Funding agencies which supports entrepreneurial development

UNIT - I No. of Hours: 09

Evolution of the concept of Entrepreneur – Characteristics – Functions and types of Entrepreneur – Entrepreneurship – Role of entrepreneurship in economic development – Women entrepreneurs – Problems of women entrepreneurs – Factors affecting entrepreneurial growth.

UNIT - II No. of Hours: 09

Mushroom cultivation: Edible mushroom – Morphology, Nutritional and medicinal value – Preparation of spawn, types of spawning – Preparation of substrate - Casing – harvesting – storage and marketing - Mushroom diseases and its management – value added products – Soup, Omlette, Samosa, Noodles, Pickles and Curry.

UNIT - III No. of Hours: 09

Composting - types of composting – aerobic and anaerobic, Drilospheres – Biology and ecological classification of earthworm – Physical and chemical effects of earthworm on soil, Vermicomposting - species employed, methods and types of production – preparation of vermiwash – Field application and crop response, Storage and marketing of composts.

UNIT - IV No. of Hours: 09

Biofertilizer – Rhizobium, BGA, Azolla, VAM – bioinoculum, mass production, field application and crop response – Biopesticide – bacteria and fungi. Production of SCP – *Spirulina* and Yeast – Herbal sale and marketing.

UNIT - V No. of Hours: 09

Finance to Entrepreneurs – Commercial banks, funding agencies – TNSCST, UGC, DST, ICMR, CSIR, and DBT. Project proposal writing – selection, formulation and financial plan - Project report preparation and submission.

#### **Text Books**

- 1. Khanka S.S (2003). **Entrepreneurial development**. 3<sup>rd</sup> edition. S.Chand & Company, New Delhi.
- 2. Kanniyan.S and Ramaswamy K (1980). **A Handbook of Edible Mushrooms**. Today's and Tomorrow's Printers, New Delhi.
- 3. Kale Radha D (1998). **Earthworm: Cinderella of organic farming**. Prism Books Pvt. Ltd., Bangalore.
- 4. Subba Rao, N.S. (1993). **Biofertilizers in Agriculture and Forestry**. 3<sup>rd</sup> edition. Oxford and IBH publication Co. Pvt. Ltd., New Delhi.

#### **Reference Books**

- 1. Shukla M.B (2007). **Entrepreneurship and small business management.** 7<sup>th</sup> edition. Kitab Mahal publication, Allahabad.
- 2. Vasant Desai (2001). **Dynamics of Entrepreneurial Development and Management.** 4<sup>th</sup> edition. Himalaya Publishing House, New Delhi.
- 3. Chang S.T and Hayes W.A (1978). **Biology and cultivation of mushrooms**. Academic Press, New York.

#### Web sources

https://www.biospace.com/article/microbiology-a-field-ripe-for-entrepreneurship/

https://extension.psu.edu/six-steps-to-mushroom-farming

https://www.systemekofungi.com/wp-content/uploads/Mushroom-Cultivation-Manual.pdf

http://www.amm-mcrc.org/publications/Biofertilizers.pdf

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	<b>✓</b>	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Fifth Semester Microbiology

## ENTREPRENEURSHIP IN MICROBIOLOGY

<b>Time: Three Hours</b>			Maximu	ım Mark: 75
	PART – A	<b>A</b> $(20 \text{ x } 1= 20 \text{ Marks})$		
	Answ	er ALL questions		
		ons carry equal marks		
-	al behaviours includes:	( ) T. 1.	9.95	(1) 411 6 1
	elems (b) Taking initiative	es (c) Taking re	esponsibility	(d) All of above
	attributes includes:	a (a) Datamai		(d) All of charre
(a) Preservence	(b) Hard working s skills includes		nation	(d) All of above
(a) Creative prob			gotiation	(d) All of above
	repreneurs was applied to but	· , ,	_	` '
century	epicheurs was applied to be	isiness initially by the i	Tellen ceollon	inst in the 16
(a) Cantillon	(b) Jan Tinbergen	(c) J.S.Mill	(b)	None of above
` '	ess function do experts agree	` /	` '	
business?	as immedian do emperos agre-	o, y o w shio wiw 10 cms o h	mor when proj	y
	(b) Marketing vision	(c) Operation	(ď	) None of above
	following is not something t			
(a) Energy	(b) Experties	(c) Money	(0	d) Time
7. From the follo	wing which one is factor at	fecting entrepreneurial	growth	
(a) Social	(b) Economic	(c) Psychologica	al	(d) All of above
	eed of entrepreneurship?			
	ovation (b) To fill gap in g		-	
	the following is the proces	s of entrepreneur devel	oping new pro	ducts that over
	nt products obsolete?			
	s model (b) Anatomization			(d) None of above
	which an individual is view			(1) A 1 !
	tus (b) Qualification			(d) Achievement
	e following shows the proce			
a) Business mod	of the following gives sugge	c) Creative flexibilit		
products?	of the following gives sugge	stions for new product	and also help	to market new
1	acts and services	b) Federal governme	ant	
c) Distribution C		d) Consumers Ques		
	following is used by entrep			nternational
	aking a major commitment		or reflect in all li	inoi nanonai
a) Merger		c) Joint venture	d) Maiority i	nterest Question

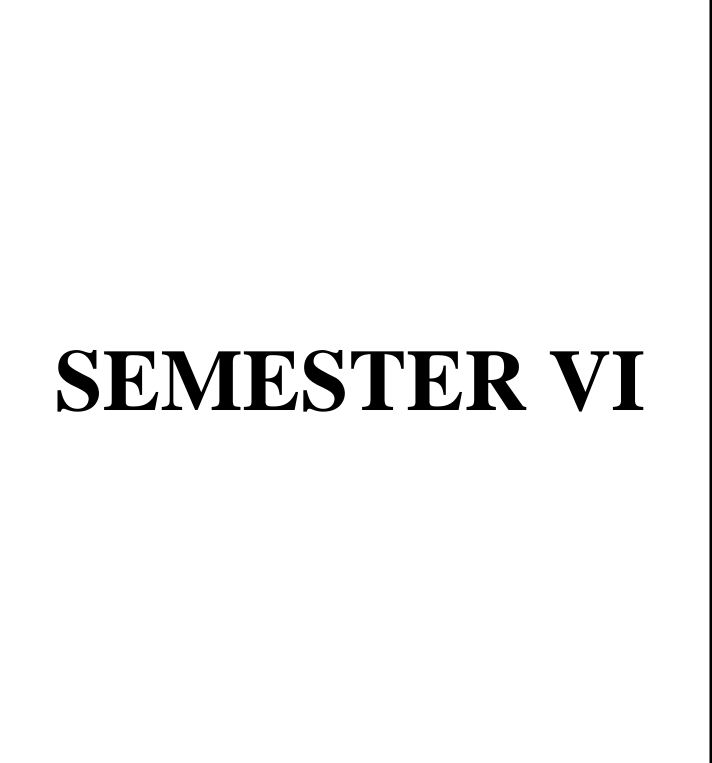
- 14. GATT is established in 1947, under:
- a) German leadership b) U.S. leadership c) French leadership d) U.K. leadership Question
- 15. The entrepreneur was distinguished from capital provider in:
- a) Middle ages
- b) 17th century
- c) 18th century
- d) 19th and 20th century
- 16. A person who managed large project was termed as the entrepreneur in the \_
- a) Earliest period
- b) Middle ages
- c) 17th century
- d) 19th and 20th century
- 17. What is the process by which individuals pursue opportunities without regard to resources they currently control?
- a) Startup management b) Entrepreneurship c) Financial analysis d) Feasibility planning
- 18. Having less than 50 percent of equity share in an international venture is called:
- a) Joint Venture b) Majority interest c) Minority interest d) Exporting
- 19. Having more than 50% ownership position that provides the entrepreneur with managerial control is called:
- a) Joint venture b) Majority interest c) Horizontal merger d) Diversified activity merger
- 20. Which one of the following is the process of entrepreneurs developing newproducts that over time make current products obsolete?
- a) New business model b) Anatomization c) None of the given options d) Creative destruction

**PART – B** (5 x 5= 25 Marks) Answer **ALL** questions All questions carry equal marks

- 21. a) Explain the factors affecting of entrepreneurial growth (or)
  - b) Describe various types of entrepreneurs.
- 22. a) Give a short notes on Mushroom diseases and its management (or)
  - b) Write about the method of spawn production.
- 23. a) Explain aerobic composting. Add notes on its uses (or)
  - b) Digramatticaly explain the structure of an earthworm.
- 24. a) Write short note on VAM bioinoculum (or)
  - b) Explain Spirulina cultivation.
- 25. a) Write about role of DST in Entrepreneurship development (or)
  - b) Describe various funding agencies.

**PART** – C (5 x 5= 25 Marks) Answer **ANY THREE** questions All questions carry equal marks

- 26. Write an essay on role of Women entrepreneurs for our national economy.
- 27. Elaborate various methods of Mushroom cultivation.
- 28. Write an essay on Vermicomposting.
- 29. Discuss in detail about the Rhizobium biofertlizer production.
- 30. Give a detailed significance of project report and project appraisal.



SEMESTER – VI
18U6MBC08
Total number of Hours: 60
Credits: 6
6 Hours/Week

## MEDICAL VIROLOGY AND PARASITOLOGY

## **Course Objectives**

To gain basic knowledge on medical virology and parasitology

To get exposure with medically important microbes and their diseases

To get expertise in diagnostic methods

To get an updated knowledge on microbes, disease control, treatment and prevention

#### **Course Outcome:**

CO1	Introduction and background on medical virology & parasitology
CO2	Able to gain knowledge on medically important common viruses
CO3	Recently emerged viral infections
CO4	Clinically important Protozoas
CO5	Clinical importance of helminthic infections

UNIT - I No. of Hours: 12

General characteristics of viruses - Viral multiplication - Cultivation of viruses - viral assay - Classification of viruses - Viroids - Prions - Antiviral agents. HSV type 1 - type 2 - type 3 - type 4 - type 5, Variola-vaccinia virus.

UNIT - II No. of Hours: 10

Rabies virus, Polio virus, Human Immunodeficiency virus, Hepatitis virus, Influenza, Mumps, measles, German measles.

UNIT - III No. of Hours: 14

**Arthropod borne and Rodent borne diseases:** Chickenguenia, Yellow fever virus, Dengue, Kyasanur forest diseases, Ebola Marbek, Ziga, Nipah virus.

UNIT - IV No. of Hours: 12

**Introduction to medical parasitology:** Classification - Common diagnostic methods in parasitology - Examination of faeces for ova and cyst - Concentration methods - Blood smear examination of parasites. *Entamoeba histolytica - Giardia lamblia - Trichomonas vaginalis - Leishmania donovani - Trypanosoma brucei - Plasmodium falciparum - Plasmodium malariae.* 

UNIT - V No. of Hours: 12

General Characteristics, life cycle, diagnosis, prophylaxis and control of Ascaris lumbricoides - Ancylostoma duodenale - Schistosoma haematobium - Taenia solium - Taenia saginata - Diphyllobothrium latum - Enterobius vermicularis- Trichuris trichiura - Wuchereria bancrofti

### **Suggested Reading**

- 1. Dimmock, NJ, Easton, AL, Leppard, KN (2007). Introduction to Modern Virology. 6<sup>th</sup> edition, Blackwell Publishing Ltd.
- 2. Carter J and Saunders V (2007). Virology: Principles and Applications. John Wiley and Sons.
- 3. Flint SJ, Enquist, LW, Krug, RM, Racaniello, VR, Skalka, AM (2004). Principles of Virology, Molecular biology, Pathogenesis and Control. 2<sup>nd</sup> edition. ASM press Washington DC.
- 4. Levy JA, Conrat HF, Owens RA. (2000). Virology. 3<sup>rd</sup> edition. Prentice Hall publication, New Jersey.
- 5. Wagner EK, Hewlett MJ. (2004). Basic Virology. 2<sup>nd</sup> edition. Blackwell Publishing.
- 6. Mathews. (2004). Plant Virology. Hull R. Academic Press, New York.
- 7. Nayudu MV. (2008). Plant Viruses. Tata McGraw Hill, India.
- 8. Bos L. (1999) Plant viruses-A text book of plant virology by. Backhuys Publishers.
- 9. Versteeg J. (1985). A Color Atlas of Virology. Wolfe Medical Publication.
- 10. Parija S.C. (2013) **Text book of Medical Parasitology.** 4<sup>th</sup> edition. All India Publishers and Distributors, New Delhi.
- 11. Chatterjee (1986). **Medical Parasitology**. Tata McGraw Hill, New Delhi.
- 12. Jagdish Chander (2012). **Text book of Medical Mycology**. 3<sup>rd</sup> edition. Mehta Publishers, New Delhi.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓		✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓		✓
CO5		✓	✓		✓

(For the candidates admitted from 2017- 18 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Sixth Semester Microbiology

#### MEDICAL VIROLOGY AND PARASITOLOGY

MEDICAL VIROLOGY AND PARASITOLOGY	
Time: Three Hours Maximum Mark: 75	
$PART - A (20 \times 1 = 20 \text{ Marks})$	
Answer ALL questions	
All questions carry equal marks	
• • •	
1. Protein coat of virus enclosing nucleic acid is called	
a. Vector b. Capsid c. Envelop d. Spikes	
2. Virus inoculation onto CAM of embryonated egg and growth identified by	
a. Pustule formation b. Edema c. Pock formation d. All	
3. Which of the following is smallest virus?	
a. HSV b. HIV c. POX virus d. Polio virus	
4. Most common and existing way to control herpes virus infections are	
a. Vaccines b. Antiviral drugs c. Immunoglobulins d. Interferons	
5. Which unique form does the rabies virus take?	
a. The virion has a dumbbell appearance b. It is shaped like a bullet from a gun	
c. The virus is star shaped d. The virion is very pleomorphic	
6. Which of the following virus contains hemagglutinin spikes?	
a. Entero virus b. Influenza virus c. VZV d. HSV	
7. Which of the following virus is arthropod born virus?	
a. HIV b. HSV c. Dengue d. Hepatitis	
8. When was smallpox eradicated from the world?	
a. In 1977 following a WHO  b. In 2000 campaign	
c. Is not yet eradicated d. In 1796 after Jenner's first vaccine	
9. Which one of the following virus can cause severe hemorrhages?	
a. Chikungunya b. Ebola c. Dengue d. Nipah	
10. Describe the morphology of a togavirus	
a. Non-enveloped with an icosahedral structure	
b. Enveloped spherical particles with an icosahedral structure	
c. Small round viruses	
d. Filamentous virus with protruding glycoproteins	
11. Which of the following is <b>not</b> a mosquito-borne illness?	
a. Nipah virus b. Zika virus c. Dengue virus d. Chikungunya	
12. Viral vaccination was invented by	
a. Jenner b. Pasteur c. Watson d. Crick	
13. The cytoplasm of the trophozoite may contain ingested when it is invasive in tissue	
a. WBCs b. Carbohydrates c. RBCs d. Macrophages	
14. The stool is the specimen for the diagnosis of the infection cause by a. Balantidium coli b. Acanthamoeba polyphaga c. Naegleria fowleri d. Leishmania donavan	ni
15. Which one of the following causes the more severe type of Malaria?	ıı
· · · · · · · · · · · · · · · · · · ·	ıv
a. Plasmodium falciparum b.Plasmodium ovale c. Plasmodium malariae d. Plasmodium viva	$\boldsymbol{x}$

16. Leishmania infection occurs due to bite of female sand fly and deposites.......

- a. Amastigote b. Promastigote c. Cyst d. Larvae
- 17. The usual infective stage of Trematodes to man is the......
- a. Cercariae b. Metacercariae c. Egg d. Miracidium
- 18. What parasite whose migrating larvae break the pulmonary capillaries of man?
- a. Ancylostoma braziliense b. Enterobius vermicularis c. Ascaris lumbricoides d. Trichuris trichiura
- 19. What parasite is associated with pork?
- a. Diphyllobothrium latum b. Taenia saginata c. Dipylidium caninum d. Taenia solium
- 20. Wuchereria bancrofti causes.....
- a. Kala-azar b. Sleeping sickness c. Lymphatic filariasis d. Black water fever

## **PART – B** (5 x 5 = 25 Marks) Answer **ALL** questions All questions carry equal marks

- 21. a. Give introduction to interferons and explain types, impact on viral infection (OR)
  - b. Explain chickenpox and shingles caused by VZV
- 22. a. Write a short note on on poliomyelitis (OR)
  - b. Explain on the causative agent of measles and pathogenesis, symptoms.
- 23. a. Summaries overall impact of *Ebola virus* infection (OR)
  - b. Explain incidence of Kyasanur forest disease and its impact
- 24. a. Explain on Giardiasis (OR)
  - b. Write on African Trypanosomiasis
- 25. a. Give account on Taeniasis (OR)
  - b. Write mode of transmission and life cycle of Ankylostoma duodenale

## **PART** – **C** (3 X 10 = 30 Marks) Answer **ANY THREE** questions All questions carry equal marks

- 26. Give introduction to antiviral agents and explain each type.
- 27. Write characteristics, pathogenesis, clinical manifestation of rabies virus.
- 28. Write infection, pathogenesis, clinical manifestation and lab diagnosis of Dengue virus.
- 29. Write in detail on acquisition of infection by *Entamoeba histolytica* and explain intestinal and extraintestinal infections and treatment.
- 30. Explain in detail on pathogenesis, clinical manifestation, lab diagnosis and treatment of *Schistosoma haematobium*.

SEMESTER – VI

18U6MBC09 Total number of Hours: 60
Credits: 5 5 Hours/Week

#### SOIL AND ENVIRONMENTAL MICROBIOLOGY

## **Course Objectives**

- To study the physico-chemical and microbiological properties of soil.
- To gain knowledge about the biogeochemical cycles and biofertilizer.
- To impart knowledge on microbial interactions in plants and animals and plant pathology.
- To understand the microbiology of air and water.
- To study the microbiology of sewage and sewage treatment methods.

### **Course Outcome:**

CO1	Able to understand soil microbiota
CO2	Concepts of metabolic pathways by soil microbes and their role
CO3	Symbiotic relationship between microbes and plants
CO4	Water quality parameters -Physico chemo parameters
CO5	They could able to perform experiments to test the quality of samples

UNIT - I No. of Hours: 12

**Introduction to soil microbiology:** Physical and chemical properties of soil - Types and significance of soil microbes – Bacteria, Fungi, Actinomycetes, Protozoa, Nematodes and Viruses. Factors affecting soil microbial population.

UNIT - II No. of Hours: 12

**Biogeochemical cycles:** Carbon, nitrogen, phosphorous and sulphur - Mechanism of nitrogen fixation - Biofertilizer – Rhizobium, Azotobacter and Cyanobacteria – Mass cultivation, field study and its applications.

UNIT - III No. of Hours: 12

**Microbial interactions and plant pathology:** neutralism, commensalism, synergism, mutualism and parasitism. Interaction of microbes with plants – Rhizosphere, Phyllosphere and Mycorrhizae. Microbe-animal interaction - Microbes in ruminants. Plant Pathology – symptoms, disease cycle and its control measures - Bacterial - Citrus canker, Fungal - Wilt of Cotton and Tikka leaf spot of groundnut, Viral – TMV.

UNIT IV No. of Hours: 12

**Microbiology of air & water** – Enumeration of bacteria from air – Air sampling devices – Air sanitation. Assessment of drinking water quality – water standards - indicator organisms – water purification – Waterborne diseases and their control measures.

UNIT V No. of Hours: 12

**Solid Waste management:** Sources and types of solid waste, Methods of solid waste disposal (composting and sanitary landfill). **Liquid waste management:** Composition and strength of sewage (BOD and COD), Primary, secondary (oxidation ponds, trickling filter, activated sludge process and septic tank) and tertiary sewage treatment. Biodegradation, Bioremediation, Biodetoriation and xenobiotics.

#### **Text Books**

- 1. Mishra R.R (2004). **Soil Microbiology**. CBS Publishers & Distributers, New Delhi.
- 2. Subba Rao (1999). **Soil Microbiology.** 4<sup>th</sup> edition. Oxford and IBH publishing Co (P) Ltd, New Delhi.
- 3. Joseph C Daniel (1999). **Environmental aspects of Microbiology**. 2<sup>nd</sup> edition. Bright Sun Publications, Chennai.
- 4. Atlas RM and Bartha R. (2000). Microbial Ecology: Fundamentals & Applications. 4<sup>th</sup> edition. Benjamin/Cummings Science Publishing, USA
- 5. Maier RM, Pepper IL and Gerba CP. (2009). Environmental Microbiology. 2<sup>nd</sup> edition, Academic Press.

#### **Reference Books**

- 1. Rangaswami.G and Bagyaraj D.J. (2009). **Agricultural Microbiology**.2<sup>nd</sup> edition. PHI Learning Pvt. Ltd., New Delhi.
- 2. Ralph Mitchell and Ji Dong Gu (2010). **Environmental Microbiology**. 2<sup>nd</sup> edition, Wiley-Blackwell, New Jersy.
- 3. Coyne MS. (2001). Soil Microbiology: An Exploratory Approach. Delmar Thomson Learning.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	<b>✓</b>	✓	✓	✓	<b>✓</b>
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	<b>✓</b>	✓	✓	✓	<b>✓</b>
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Sixth Semester Microbiology

## SOIL AND ENVIRONMENTAL MICROBIOLOGY

Time: Three Hours	Maximum Mark: 75
PART - A (20  x  1 = 20  Marks)	
Answer <b>ALL</b> questions	
All questions carry equal marks	
<ol> <li>The population of algae in soil is that of either bacteria</li> <li>Generally smaller than b. Generally greater than c. Equal to</li> <li>Soil organic matter a good indicator of</li> <li>Biological health b. Chemical health c. Physical health</li> <li>Soil microorganisms are most active at</li> </ol>	d. None of the above
a. 15-20°C b. 20-25°C c. 34-36°C d. 40-45°C	
<ul> <li>4 play a key role in the transformation of rock to soil</li> <li>a. Cyanobacteia b. Pectin decomposing bacteria</li> <li>c. Nitrifying bacteria d. De-nitrifying bacteria</li> <li>5. The association which involves the exchange of nutrients between</li> </ul>	en two species is referred to as
a. Mutualism b. Syntrophism c. Commensalism d. 6. Which of the following conditions decreases the level of denitrifia. Abundance of organic matter b. Acidic pH c. Elevated tempera oxygen	ication?
7. Which of the following is a symbiotic nitrogen fixing bacteria? a. <i>Rhizobium trifolii</i> b. <i>Clostridium pasteurianum</i> c. <i>Azotobac</i> 8. The word Rhizosphere and Phyllosphere is respectively given by	,
a. Hiltner and Ruinen b. Ruinen and Hiltner c. Winogradsky an	d Beijernickia d. Frank
9. Tikka disease of groundnut is caused by	F
a. <i>Puccini</i> b. <i>Aspergillus</i> c. <i>Cercospora</i> d. <i>I</i> 10. Ammonia oxidizers and nitrite oxidizers are	
a) Gram-negative chemolithotrophs b) Gram-positive cl	hemolithotrophs
a) Gram-negative chemolithotrophs c) Gram-negative photolithotrophs d) Gram-positive pl	hotolithotrophs
11. Which among the following is not an ammonia-oxidizing bacter	ia?
<ul><li>a) Nitrosomonas europaea</li><li>b) Nitrosovibrio tenuis</li><li>c) Nitrospina gracilis</li><li>d) Nitrosococcus oceanus</li></ul>	
12. How much time does nitrifying bacteria requires to grow at an i	ncubation of 250 to 300 C?
	4 months
13. Which of the following is/are inorganic gas(es)?	
a. Carbon monoxide b. Hydrogen sulphide c. Chlorine	d. All of the above
14. Which type of Hepatitis spreads through polluted water?	J. Darle A. and D.
a. Hepatitis A b. Hepatitis B c. Hepatitis C	d. Both A and B
15. Haemophillia' is a disease associated with	

- 16. The following Disease is water borne
- a. Typhoid
- b. Tuberculosis
- c. Hepatitis B
- d. Scurvy
- 17. The biological oxygen demand (BOD) would be most directly affected by the presence of which of the following pollutants?
- a. Heavy metals
- b. Organic wastes
- c. Salt (Sodium chloride)
- d. Waste minerals from mining e. Fertilizer runoff from farms
- 18. The microorganisms that is mainly used as an indicator of fecal pollution in water is:
- a. Escherichia coli
- b. Clostridium tetani
- c. Clostridium botulinum

- d. Cyanobacteria
- e. All of these
- 19. Which of the following waste water treatments is most likely to produce carcinogens as a byproduct?
- a. Chlorination
- b. Ozonation
- c. Ultraviolet light (UV)

- d. Sand filtration
- e. Carbon filtration
- 20. A major disadvantage of bioremediation is: a. Long times may be required b. It is n
  - b. It is more expensive than other treatments
- c. It can not be used to treat contamination with hydrocarbons
- d. It requires removing contaminated soil to a bioreactor
- e. It requires introducing new microorganisms into an environment

## $PART - B (5 \times 5 = 25 \text{ Marks})$

Answer **ALL** questions

All questions carry equal marks

- 21. a) Write about properties of soil (OR)
  - b) Explain the role of Bacteria and Actinomycetes in soil?
- 22. a) Write short notes on ammonification, nitrification and denitrification? (OR)
  - b) Types of Biofertilizers with examples.
- 23. a) Explain the types of microbial interaction? (OR)
  - b) Write a short note on Mycorrhizae
- 24. a) How to Enumerate bacteria from air? (OR)
  - b) Types of Air sanitation methods?
- 25. a) Role of microbes in Sewage treatment (OR)
  - b) Effect of thermal pollution in the environment?

## $PART - C (3 \times 10 = 30 \text{ Marks})$

Answer **ANY THREE** questions

All questions carry equal marks

- 26. Explain the significance of soil microbes: Fungi, Microalgae, Protozoa and Viruses?
- 27. Write an essay on Nitrogen cycle.
- 28. Explain the bacterial diseases & fungal disease in plant?
- 29. Write in detail about the waterborne diseases?
- 30. Briefly explain the Sewage treatment?

SEMESTER – VI
18U6MBC10
Credits: 5
Credits: 5

CORE - X
Total number of Hours: 60
5 Hours/Week

FOOD AND DAIRY MICROBIOLOGY

## **Course Objectives**

- To gain knowledge about the microorganisms involved in food
- To impart the idea in food spoilage
- To gain the knowledge in food preservation.
- To study the food borne infections
- To study the rules and regulations of food sanitation

#### **Course Outcome:**

CO1	Food pathogens and their Phsico-chemico parameter analysis
CO2	Spoilage of food by various microbes
CO3	Food Preservation methods
CO4	Able to understand microbial Fermented products
CO5	Food intoxication and determination of food pathogens

UNIT - I No. of Hours: 12

Foods as a substrate for microorganisms – Importance of microorganisms in food - Bacteria, Mold and Yeasts. Sources of food contamination. Factors affecting the Growth - Intrinsic factors - (pH, moisture, oxidation - reduction potential, and nutrient content), extrinsic factors - (temperature, relative humidity, gases and microbial activities) and inhibitory substances.

UNIT - II No. of Hours: 12

Microbial spoilage of various foods - General Principles underlying food spoilage and contamination - Spoilage and preservation of vegetables and fruits, meat and eggs, dairy products and sea foods.

UNIT - III No. of Hours: 12

Principles and methods of food preservation – Physical and chemical methods - Physical methods – Asepsis, temperature (low, high, canning, drying), irradiation, hydrostatic pressure, high voltage pulse, microwave processing and aseptic packaging – tetra packing. Chemical methods - salt, sugar, organic acids, SO<sub>2</sub>, nitrite and nitrates, ethylene oxide, antibiotics and bacteriocins.

UNIT - IV No. of Hours: 12

Fermented food products - Dairy starter cultures, fermented dairy products - yogurt, acidophilus milk, kumiss, kefir, dahi and cheese. Other fermented foods - dosa, sauerkraut, soy sauce and tampeh. Probiotics - Health benefits, types of microorganisms used, probiotic foods available in market, GRAS (General Regard as Safe).

UNIT - V No. of Hours: 12

Food borne diseases and Food sanitation: Food intoxications and food borne diseases. Rapid detection methods of food borne pathogens. Food sanitation and control.

## **Text Books**

- 1. Vijaya Ramesh K (2007). Food Microbiology. First edition, MJP Publishers, Chennai.
- 2. Adams MR Moss MO (2004). Food Microbiology, 2<sup>nd</sup> Edition, Panima Publishing House, New Delhi.
- 3. James M Jay (2003). Modern Food Microbiology. 4<sup>th</sup> Edition, CBS Publishers & Distributors, New Delhi

### **Reference Books**

- 1. Frazier WC and Westhoff DC (1988). Food Microbiology, 4<sup>th</sup> Edition, Mc Graw Hill, New York
- 2. Banwart JM. (1987). Basic Food Microbiology. 1<sup>st</sup> edition. CBS Publishers and Distributors, Delhi, India.
- 3. Jay JM, Loessner MJ and Golden DA. (2005). Modern Food Microbiology. 7<sup>th</sup> edition, CBS Publishers and Distributors, Delhi, India.
- 4. Sivashankar B Moss (2011). Food Processing and Preservation. Eighth edition, PHI Learning P.Ltd., New Delhi.
- 5. Roday, S. (1998). Food Hygiene and Sanitation. Tata Mcgraw Hill Publications.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	<b>✓</b>			✓	✓
CO2	✓	✓		✓	✓
CO3	✓	✓	✓	✓	
CO4	✓	✓	✓	✓	
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

## **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Sixth Semester Microbiology

## FOOD AND DAIRY MICROBIOLOGY

Time: Three Hours  Maximum Mark: 75								
$\mathbf{PART} - \mathbf{A} (20 \times 1 = 20 \text{ Marks})$								
Answer ALL questions								
All questions carry equal marks								
• • •								
1. Spoilage in food because of microbial activity can be prevented or delayed by								
a. Prohibiting the entry of micro-organisms in food b. Physical removal of micro-organisms								
c. Hindering the activity of micro-organisms d. All of above								
2. The growth of aerobic food spoilage and pathogenic microorganisms can be suppressed by								
a. Humectants b. Exhausting c. Both a and b d. None of above								
3. The target microorganism in canning is								
a. Clostridium botulinum b. Streptococcus thermophillus c. PA 3679 d. Lactobacillus								
bulgaricus								
4. Pasteurization is the heat treatment designed primarily to kill								
a. Vegetable forms of microorganisms b. All form of microorganisms c. Spore d. None of								
above								
5. In spore forming bacteria maximum resistance occurs at pH								
a. 4 b. 5 c. 6 d. 7								
6. The time required to kill microorganism at a given lethal temperature is known as								
a. Z value b. D value c. C value d. B value								
7. The microorganisms multiply and die in								
a. Geometric order b. Logarithmic order c. A-logarithmic order d. None of above								
8. Acetobactor aceti convertsinto acetic acids								
a. Ethyl alcohol b. Glucose c. Methyl alcohol d. Starch								
9. Two types of fermentations are carried out for the production of								
a. Pickle b. Yoghurt c. Vinegar d. Sausages								
10. In bread manufacturing, alcoholic fermentation is carried out by								
a. Streptococcus thermophillus b. Saccharomyces cerevisae								
c. S. carlsbergensis d. Lactobacillus bulgaricus								
11. Clostridium botulinum mainly result in spoilage of foods								
a. High acid Food b. Acidic Food c. Medium acid Food d. Low acid								
Food								
12. Any change that renders food unfit for human consumption is called								
a. Processing b. Spoilage c. Deterioration d. Preservation								
13. The temperature resistance of microorganism in high acid food is								

c. Low

a. High

b. Medium

d. No effect

14. Food intoxication is the ingestion of a. enzymes producing microorganism b. Toxin producing microorganism c. Non of both d. Both of these 15. Clostridium Botulinum is ----a. Bacteria b. Mold c. Yeast d. Virus 16. Thermophiles grows at a. 8 to 45°C b. 25 to 30°C c. 0 to 20°C d. 50-600 C 17. Type of yeast used for alcoholic fermentation is a. Saccharomyces Cerevisiae b. Streptococcus thermophillus c. Acetobacter acceti d. Clostridium botulinum 18. Lactic acid bacteria include ----a. Lactococcus lactis b. Lactococcus cremoris c. Bifidobacterium d. All above 19. Which of the following products have higher acidity and lacks aroma? a) Cultured buttermilk b) Cultured sour cream c) Bulgarian milk d) Acidophilus milk 20. The microbiological examination of coliform bacteria in foods preferably use a. MacConkey broth b. MacConkey agar c. eosine Methylene blue agar d. all of these

## **PART - B** (5 x 5 = 25 Marks)

Answer **ALL** questions

All questions carry equal marks

- 21. a) Write about microorganism involved in food spoilage (or)
  - b) Discuss the principles of food preservation.
- 22. a) Write short notes on contamination and spoilage of fruits and vegetables(or)
  - b) Explain the spoilage and preservation of fish and other sea foods.
- 23. a) Write short notes on botulism (or)
  - b) Describe the parasitic infections.
- 24. a) Discuss about the methods of fermentation (or)
  - b) Give short notes on production of beer.
- 25. a) Write short notes on yoghurt production (or)
  - b) Discuss about the microflora of milk.

## PART - C (3 X 10 = 30 Marks) Answer **ANY THREE** questions

All questions carry equal marks

- 26. Briefly explain the factors influencing microbial growth in food.
- 27. Explain the contamination, spoilage and preservation of Cereal and Cereal product.
- 28. Discuss the food borne intoxication in detail.
- 29. Give a brief account on production of Wine.
- 30. Detail about the role and responsibilities of food control agencies and its regulations.

SEMESTER – VI
18U6MBE03

Credits: 4

ELECTIVE - II
Total number of Hours: 45
4 Hours/Week

#### MICROBIAL DIAGNOSIS IN HEALTH CLINICS

## **Course Objectives**

- To gain knowledge about the microbial diseases.
- To impart knowledge on clinical sample collection.
- To gain knowledge about microbial characters in selective media.
- To study the different detection methods.
- To gain the knowledge on antimicrobial testing & MIC.

#### **Course Outcome:**

CO1	Microbial disease diagnosis methods
CO2	To understand the clinical microbiology
CO3	Able to understand the microscopic examination
CO4	Able to understand molecular identification by molecular techniques
CO5	To understand the antibiotics test

UNIT - I No. of Hours: 9

**Importance of Diagnosis of Diseases:** Bacterial, Viral, Fungal and Protozoan - Diseases of various human body systems - Disease associated clinical samples for diagnosis.

UNIT - II No. of Hours: 9

**Collection of Clinical Samples:** Guidelines for the collection — Transport - Processing and analysis of samples - oral cavity, throat, sputum, skin scrapings, Blood, CSF, urine and faeces and its precautions - Storage method of clinical samples in laboratory.

UNIT - III No. of Hours: 9

**Direct Microscopic Examination and Culture:** Examination of sample by staining - Gram stain, Ziehl-Neelson staining for tuberculosis, Giemsa -stained thin blood film for malaria. Preparation and use of various selective media - Distinct colony properties of various bacterial pathogens in selective medium.

UNIT - IV No. of Hours: 9

**Serological and Molecular and rapid detection Methods:** Serological Methods – ELISA & immunofluorescence. Molecular methods - PCR & Nucleic acid probes. Rapid Detection methods - Typhoid, Dengue and HIV using diagnostic kits.

UNIT - V No. of Hours: 9

**Testing for Antibiotic Sensitivity in Bacteria:** Importance, Determination of resistance/sensitivity of bacteria using disc diffusion method - Determination of minimal inhibitory concentration (MIC) of an antibiotic by serial double dilution method.

## **Text Books**

- Ananthanarayan R and Paniker CKJ (2009). Textbook of Microbiology, 8<sup>th</sup> edition, Universities Press Private Ltd.
- 2. Jawetz, Melnick and Adelberg's Medical Microbiology. 26<sup>th</sup> edition. McGraw Hill Publication.

## **Reference Books**

- 1. Topley & Wilsons Microbiology & Microbial Infections 9<sup>th</sup> Edition.
- 2. Brooks G.F., Carroll K.C., Butel J.S., Morse S.A. and Mietzner, T.A. (2013).
- Randhawa, VS, Mehta G and Sharma KB (2009) Practicals and Viva in Medical Microbiology 2<sup>nd</sup> edition, Elsevier India Pvt Ltd.
- 4. Tille P (2013) Bailey's and Scott's Diagnostic Microbiology, 13th edition, Mosby.

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

# **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Sixth Semester Microbiology

# MICROBIAL DIAGNOSIS IN HEALTH CLINICS

me: Three Hours			Maxii	mum Mark: 75
	PART -	$\mathbf{A}$ (20 x 1= 20 Marks)		
	Ansv	ver <b>ALL</b> questions		
	All quest	ions carry equal marks		
	•	J 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		
1. Enrichment media is alw		o Comi calid madium	d Cala	-ti di
<ul><li>a. Liquid medium</li><li>b.</li><li>2. Mycobacterium culture g</li></ul>			d. Sele	ctive medium
a. Lowenstein-Jensen med			r d M	IacConkey agar
3. What is the temperature of		igai C. Ivutilent aga	u. 1v.	iacconkcy agai
a120 degree C b.0	degree C	c150 degree C	d196 deg	ree C
a120 degree C b.0 4. Which of the following	g method can be	used to determine the	number of b	acteria quantitatively?
a. Streak plate b. Spre				
5. Which of the following a			•	• •
a. Agar slant is covered w				degree to -78 degree C
c. Vials are connected to l			s dehydrated	
6. Which of the following is	s a function of cryo	protective agents?		
a. For long-term preservat	ion of cultures	b. Prevents cell damag d. To trap the liquid ni	ge due to ice c	rystal formation
c. Prevents formation of ic	e	d. To trap the liquid ni	itrogen	
7. Crystal violet isa. Primary stain b. M.		ondary etain d	All of these	
8. The system of antiseptic			All of these	
a. John Tyndall b			d. Rol	oert Koch
9. Tuberculosis is a		C. Bould I doctor	<b>3.</b> 100	, <b>4.1 ( 1.3 ( 1.</b>
a. Water borne disease	b. Air borne diseas	se c. Food borne disea	ise d. A	Althropod disease
10. EMB agar is a				-
a. Enriched media b. D			d. Enrichn	nent media
11. Mycobacterium culture				
a. Lowenstein-Jensen med			ar d. N	AacConkey agar
12. Whose is known as Fath a. Robert Koch b. Ed		c. Louis Pasteur	d. Fle	mina
13. Nichrome loop wire is				ming
a. Pour plate b. Stre				technique
14. Which one of the follow	ving is true?			
a. Agar has nutrient proper	rties	b. Chocolate medium is s	elective medi	ım
c. Nutrient broth is basal n	nedium	d. Liquid medium is selec	ctive medium	
15. Counter stain used in gr				
a. Safranin b. C	rystal violet c	. Carbol fuschion	d. Ace	toacramine
16. Anthracis was isolated by	•			
a. Robert Koch	b. Edward Jenner	c. Anton Von Leeuwe	enhoek	d. Fleming

b. Chocolate medium is selective medium

d. Liquid medium is selective medium

17. Which one of the following is true?

a. Agar has nutrient propertiesc. Nutrient broth is basal medium

- 18. Counter stain used in gram staining is -----
- a. Safranin
- b. Crystal violet
- c. Carbol fuschion
- d. Acetoacramine

- 19. Mycobacterium culture grown on -----
- a. Lowenstein-Jensen medium
- b. blood agar
- c. Nutrient agar
- d. MacConkey agar

- 20. Whose is known as Father of Immunology?
- a. Robert Koch
- b. Edward Jenner
- c. Louis Pasteur
- d. Fleming

**PART – B** (5 x 5 = 25 Marks) Answer **ALL** questions All questions carry equal marks

- 21. a) Write a short note on bacterial diseases (OR)
  - b) Write a short note on fungal diseases.
- 22. a) Discuss about collect and transport of clinical sample (OR)
  - b) Describe the storage method of clinical sample.
- 23. a) Describe the gram staining technique (OR)
  - b) Explain the Giemsa stained thin blood film for malaria.
- 24. a) Briefly explain about typhoid (OR)
  - b) Briefly explain about Dengue.
- 25. a) Discuss about serial tube dilution method (OR)
  - b) Write about the disc diffusion method.

**PART** – C (3 X 10 = 30 Marks) Answer **ANY THREE** questions All questions carry equal marks

- 26. Describe the importance of diagnosis of diseases.
- 27. Describe the collection and transport of clinical samples.
- 28. Explain Gram staining and acid fast staining.
- 29. Discuss in detail about PCR.
- 30. Briefly explain about MIC.

SEMESTER – VI
17U6MBE04
Total number of Hours: 45
Credits: 4
4 Hours/Week

# QUALITY CONTROL IN FOOD MICROBIOLOGY

# **Course Objectives:**

- GLP practices are intended to promote the quality and validity of test data.
- To get an idea for food business sets around producing and providing safe.
- To be able to differentiate between different enumeration techniques and learn when each should be used.
- To gain knowledge on spoilage microorganisms affects the appearance, smell, texture and taste.
- To Identify sources of potential errors during production and confirm the quality of the final product

#### **Course Outcome:**

CO1	Able to understand good laboratory practices
CO2	Able to understand the importance and food safety method
CO3	To gained knowledge about microbes and their food product
CO4	Able to understand food spoilage methods
CO5	Able to understand food preservation technologies

UNIT - I Total No. of hours: 45

Good laboratory practices (GLP), Good Microbiological Practices (GMP). Quality policy, quality objectives of food processing company, Standard Operating Procedures, Work instructions, Good Handling Practices (GHP) & GMP checklist.

UNIT - II Total No. of hours: 45

Importance and significance of microorganisms in food safety - Food and Drug Administration (FDA) and its regulation - Factors affecting the growth of micro organisms in food - intrinsic (pH, moisture, oxidation-reduction potential and nutrient content) and extrinsic (Temperature, relative humidity, gases and microbial activities).

UNIT - III Total No. of hours: 45

Determination of micro organisms and their products in food: sampling, sample collection, transport and storage, sample preparation for analysis. Microscopic and culture dependent methods- direct microscopic observation, culture enumeration and isolation methods.

UNIT - IV Total No. of hours: 45

Food spoilage: characteristic features, dynamics and significance of spoilage of different groups of foods - cereal and cereal products, vegetables and fruits, meat poultry and sea foods, milk and milk products, packed and canned foods.

UNIT- V Total No. of hours: 45

Rules and regulations for setting up of a processing unit. Criteria for ingredients and finished products. Aspects of microbiological safety in food preservation technologies, Establishment and implementation of HACCP, Continuous Assessment System, Total quality management and quality audits in food industries.

#### **Suggested Books:**

- 1. Frazier, W.C. (1988) Food Microbiology, Mc Graw Hill Inc. 4<sup>th</sup> Edition.
- 2. The training manual for Food Safety Regulators. Vol.II- Food Safety regulations and food safety management. (2011) Food safety and Standards Authority of India. New Delhi.
- 3. Fundamentals of Dairy Microbiology by Prajapati.
- 4. Pelczar, M.I., and Reid, R.D. (2009) Microbiology, 5th Ed., McGraw Hill Inc., New York.
- 5. James, M.J. (2007) Modern Food Microbiology, 2nd Ed., CBS Publisher, New Delhi
- 6. Adams, M.R., and Moss, M.G., (2005) Food Microbiology, 1st Ed., New Age International (P) Ltd., New Delhi.

#### **Mapping**

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

#### **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Sixth Semester Microbiology

# QUALITY CONTROL IN FOOD MICROBIOLOGY

Time: Three Hours Maximum Mark: 75

PART - A (20 x 1 = 20 Marks)

Answer **ALL** questions

All questions carry equal marks

<b>1</b>
1. Good work practices include
<ul><li>a. smelling and tasting chemicals</li><li>b. not washing hands before and after lab</li><li>c. confining long hair and loose clothing</li><li>d. using damaged equipment and glassware.</li></ul>
c. confining long hair and loose clothing d. using damaged equipment and glassware.
2. Chemical, reagents or broth cultures should be pipetted by?
a. mouth b. pipetter c. ear d. nose
3. The desire to maintain a safe laboratory environment for all begins with?
a. prevention b. microbiology c. ubiquity d. accidents
4. To prevent the contamination of microscopes and surrounding areas disenfect/clean used slides,
prepared by student, with
a. 70% ethanol and lens paper b. acetone and lens paper
c. 5% methylene blue and lens paper d. water and lens paper
5 is needed as a source of nutrient for the growth and reproduction of microbes.
a. pathogens b. reagents c. bacteria d. media
6. The growth of aerobic food spoilage and pathogenic microorganisms can be suppressed by
a. Humectants b. Exhausting c. Both a and b d. None of above
7. Pasteurization is the heat treatment designed primarily to kill
a. Vegetable forms of microorganisms b. All form of microorganisms c. Spore d. None of
above
8. Clostridium botulinum mainly result in spoilage of foods
a. High acid Food b. Acidic Food c. Medium acid Food d. Low acid Food
9. Bacteria which is present in raw or undercooked meat, eggs, sea food and unpasteurized milk is
a. <i>E.coli</i> b. <i>Salmonella</i> c. <i>Staphylococcus</i> d. Cyanobacteria
10. Milk and curry left over can be turned into sour and spoiled at
a. high temperature b. very low temperature c. room temperature d. constant temperature
11. Diarrhea, vomiting and severe abdominal cramps shows their sign in
a. food poisoning b. constipation c. heart diseases d. muscle cramps
12. The undesirable change in a food that makes it unsafe for human consumption is referred as
a) food decay b) food spoilage c) food loss d) all of the above
13. Common food poisoning microbes are
a) Clostridium and Salmonella b) Clostridium and E.coli c) E.coli and Salmonella d) Clostridium and Streptococcus
14. Which of the following statements are true regarding Staphylococcus food poisoning is
a) an enterotoxin b) causes gastroenteritis c) is produced by <i>Staphylococcus aureus</i> d) all of
these

15. Bacterial cell grown on hydrocarbon wastes from the petroleum industry are a source of ----a) carbohydrates b) proteins c) vitamins 16. Which of the following products have higher acidity and lacks aroma? a) Cultured buttermilk b) Cultured sour cream c) Bulgarian milk d) Acidophilus milk 17. The microbiological examination of coliform bacteria in foods preferably use a. MacConkey broth b. MacConkey agar c. eosine Methylene blue agar d. all of these 18. How many HACCP reguatios are there b. 3 a. 2 c. 4 d. None of these 19. Which of the following disease is best diagnosed by serologic means? a. Pulmonary tuberculosis b. Gonorrhea c. Actinomycosis d. Q Fever 20. HACCP stands for ----a. Hazard Activity Critical Control Plan b. Hazard Analysis Critical Control Points c. Hygiene Analysis Critical Control Points d. Hygiene Analysis Contamination Control Plan

# $PART - B (5 \times 5 = 25 \text{ Marks})$

### Answer **ALL** questions

All questions carry equal marks

- 21. a. Write about the good laboratory practices (or)
  - b. Discuss about the standard operating procedures.
- 22. a. Write a short note on Food and Drug administration (or)
  - b. Describe about the factors affecting the growth of Micro organisms if food.
- 23. a. Write about the Sample preparation for analysis (or)
  - b. Describe about the direct microscopic observation of pathogens in food sample.
- 24. a. How the micro organisms spoiled the packed and canned foods (or)
  - b. Write about the factors involved in food spoilage.
- 25. a. Explain about the HACCP (or)
  - b. Write a short note on total quality management.

#### $PART - C (3 \times 10 = 30 \text{ Marks})$

#### Answer **ANY THREE** questions

All questions carry equal marks

- 26. Discuss about the quality objectives of food processing company.
- 27. Explain importance and significance of micro organisms in food safety.
- 28. Explain about the determination of organisms if food products.
- 29. Discuss the significance of organisms in spoilage of different groups of foods.
- 30. Write the Rule and regulations for setting up of a processing unit.

SEMESTER – VI
18U6MBS04

SBEC - IV
Total number of Hours: 30

Credits: 2 2 Hours/Week

#### ADVANCES IN MICROBIOLOGY

### **Course Objectives**

- To understand quorum sensing.
- To gain knowledge about metagenomics.
- To become familiar with microbial fuel cell (MFC).
- To understand biotechnological potential of algae.
- To gain knowledge about modern trends in microbial production.

#### **Course Outcome:**

CO1	Able to understand the quorum sensing and their applications
CO2	Able to understand the human metagenomics projects
CO3	To understand the Microbial fuel cell Technology
CO4	Able to understand the animal cell culture methods
CO5	To understand the Modern trends in microbial production

UNIT – I No. of Hours: 06

**Quorum sensing:** Virulence factors associated with quorum sensing - molecular mechanisms. Quorum quenching - prokaryotic to prokaryotic quorum quenching - Eukaryotic to prokaryotic quorum quenching - applications of quorum quenching.

UNIT – II No. of Hours: 06

**Metagenomics:** History and development - Steps involved and application of metagenomics - bacterial diversity using metagenomics approach - Prospecting genes of biotechnological importance using metagenomics - Basic knowledge of metatranscriptomics, metaproteomics and metabolomics.

UNIT - III No. of Hours: 06

**Microbial fuel cell (MFC) Technology:** Microorganisms involved in MFC - Working principle - Interaction between microbes and electrodes - Design and Architecture of MFC - Types: Single chambered, double chambered. Application of MFC in Bio-hydrogen production, waste water treatment.

UNIT - IV No. of Hours: 06

**Animal Cell Culture Technology:** Introduction – types of cells - cell culture media and supplements, adherent cells – Vero, Hep-2, HepG-2, HeLa, MDCK, BHK – cultivation - subculturing – preservation – revival.

UNIT – V No. of Hours: 06

**Modern trends in microbial production:** Microbial production of bioplastics, Bioinsecticide (thruricide), biopolymer (dextran, alginate, Xanthan), Biofertilizer -  $N_2$  fixer - Azotobacter, phosphate solubilizer, Single cell protein (SCP).

#### **Text Books**

- 1. Purohit SS (2005). **Biotechnology: Fundamentals and Applications.** 3<sup>rd</sup> Edition Agrobios (India).
- 2. Sathyanarayana U (2005). **Biotechnology.** 1<sup>st</sup> Edition, Books and Allied (P) Ltd., Kolkata.
- 3. Dubey RC (2006). **A Text Book of Biotechnology.** 4<sup>th</sup> Edition. S.Chand & Company (P) Ltd., New Delhi.
- 4. Jogdand SN (2010). Environmental Biotechnology. Himalaya Publishing House, New Delhi.

#### **Reference Books**

- 1. Bernad R Glick (2010). **Molecular Biotechnology Principles and Applications of Recombinant DNA**. 4<sup>th</sup> Edition, ASM Press, Washington, D.C.
- 2. Maheswari DK and Dubey RC (2008). **Potential Microorganisms for Sustainable Agriculture**. I K International Publishing House Pvt. Ltd.
- 3. Sahoo D and Kaushik BD (2012). **Algal Biotechnology and Environment**.1<sup>st</sup> Edition, I K International Publishing House Pvt. Ltd.
- 4. Thatoi HN and Mishra BB (2011). **Microbial Biotechnology: Methods and Applications.** 1<sup>st</sup> Edition, Alpha Science International Ltd.
- 5. Fraser CM, Read TD and Nelson KE. (2004). Microbial Genomes. Humana Press.
- 6. Madigan MT, Martink JM, Dunlap PV and Clark DP (2014). **Brook's Biology of Microorganisms**, 14<sup>th</sup> edition, Pearson-Bejamin Cummings.

#### **Mapping**

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓

(For the candidates admitted from 2017- 18 onwards)

# **B.Sc., DEGREE EXAMINATIONS**

----- / ----- 2018.

Sixth Semester Microbiology

# ADVANCES IN MICROBIOLOGY

Time: Three Hours	Maximum Mark: 75

PART - A (20 x 1 = 20 Marks)

Answer <b>ALL</b> questions
All questions carry equal marks
1. Form of gene expression which is regulated in response to cell density. Used as form of communication between cells through autoinducers. Commonly found in biofilmsa. Signal peptides b. Quorum sensing c. Autoinducer d. Biofilm 2. Which of the following mode of microbial communication is likely to be faster, and more effective for less denser populations?
<ul><li>a. Quorum sensing making use of chemical signals</li><li>b. Both will happen at almost same speed</li><li>c. Communication using physical signals such as sound</li><li>d. None of the above</li></ul>
3. Signal molecule fits the binding site on its complementary receptor called as
a. Specificity b. Amplification c. Integration d. Cooperativity
4. The information which is represented by a signal is detected by specific receptors and converted
to a cellular response; this conversion is called
a. Signal amplification b. Signal transversion c. Signal transduction d. Signal integration
5. Restriction fragment length polymorphisms (RFLPs)
a. Are used to determine the position of restriction sites in a genome
b. Are used in physical mapping
c. Are used in genetic mappind. Usually occur as multiple (more than 2) alleles in a genome
6. Which of these projects would be best suited for Next Generation Sequencing?
<ul><li>a) To determine if a tumour sample contains a common missense mutation</li><li>b) To find the transcriptome of a tumour sample</li></ul>
c) To genotype ten genomic DNA samples for a known single nucleotide polymorph
d) All of the above
7. What is metagenomics?
a. Genomics as applied to a species that most typifies the average phenotype of its genus
b. The sequence of one or two representative genes from several species
c. The sequencing of only the most highly conserved genes in a lineage
d. Sequencing DNA from a group of species from the same ecosystem
8. What is proteomics?
a. The linkage of each gene to a particular protein
b. The study of the full protein set encoded by a genome
c. The totality of the functional possibilities of a single protein
d. The study of how amino acids are ordered in a protein
e. The study of how a single gene activates many proteins
9. A fuel cell is used to convert chemical energy into
a. Mechanical energy b. Solar energy c. Electrical energy d. Potential energy
10 and suitable catalyst are required to promote high rate of electrode processes.

a. Lower temperature b. Higher temperature c. Moderate temperature d. Very low temperature

11is the device used to measure the emf of the cell.					
a. Voltmeter b. Potentiometer c. Ammeter d. Multimeter					
12. The temperature maintained in the standard hydrogen electrode is					
a. 22°C b. 23°C c. 24°C d. 25°C					
13. Hybridoma cells have an application to produce					
a. Antigens b. Antibodies c. Cancer cells d. Cell lines					
14. The following are a list of essential components of cell culture media. Match them to the					
requirements for effective cell culture which they fulfil?					
a. Phenol red b. Glutamine c. Inorganic salts d. Bicarbonate					
15. Eicosanoids is a type of					
a. Hormone b. Antibiotic c. Vaccine d. Antigen					
16. The first vaccine developed from animal cell culture was					
a. Hepatitis B vaccine b. Înfluenza vaccine c. Small Pox vaccine d. Polio vaccine					
17. Biofertilizers include					
a. Cow dung manure and farmyard waste b. Quick growing crop ploughed back					
c. BGA/Anabaena and Azolla d. All of the above					
18. Which of the following material is used as bioplastic?					
a. Polystyrene b. Polypropylene c. Polyhydroxybutrate d. Dextran					
19. Technique of SCP is introduced by					
a. Gregor Mendel b. Louis Pasteur c. Professor Scrimshaw d. Ian Wilmot					
20. By using single-cell protein, amount of protein that can be produced by algae grown in ponds					
(per acre) is					
a. 20 tons b. 30 tons c. 40 tons d. 50 tons					
<b>DADT D</b> (5 v 5 – 25 Morles)					
$\mathbf{PART} - \mathbf{B} \text{ (5 x 5 = 25 Marks)}$ Answer <b>ALL</b> questions					
All questions carry equal marks					
21. a) Describe the molecular mechanism of quorum sensing in Myxobacteria (OR)					
b) Shortly explain about the application of biofilm					
22. a) Write about the definition and types of biofilm (OR)					
b) Discuss the impact factor of biofuel production					
23. a) Write the short notes on metal recovery of copper and iron (OR)					
b) Write the brief account on the biodegradable plastics.					
24. a) Discuss about Rhizosphere, Rhizoplane & Phyllosphere (OR)					
b) Write short note on Azospirillum					
25. a) Explain the role of microalgae as colourant (OR)					
b) Describe the Spirullina cultivation method in detail.					
o) Beserve the spirannia easi variou memoa in actain					
$PART - C (3 \times 10 = 30 \text{ Marks})$					
Answer ANY THREE questions					
All questions carry equal marks					
26. Discuss about the bacterial quorum sensing.					
27. Write the essay notes on bioenergy production.					

28. Discuss about the processing of microbial leaching. 29. Describe types and application of Biopesticide.

30. Give a detailed account on Mass cultivation of Rhizobium.

SEMESTER – VI
17U6MBCP06
Credits: 3
CORE - VI
Total number of Hours: 60
6 Hours/Week

# MAJOR PRACTICAL VI – MEDICAL VIROLOGY AND PARASITOLOGY, SOIL AND ENVIRONMENTAL MICROBIOLOGY, FOOD AND DAIRY MICROBIOLGY

# **Course Objectives:**

- To obtain knowledge about virus identification methods
- To gain information about the identification of human parasites
- To know the techniques in the isolation of bacteria from root nodules
- To update the identification methods used in assess the water quality
- To get knowledge about the microbes from spoiled food materials

#### **Course Outcome:**

CO1	To understand the hemagglutination techniques
CO2	Able to understand the cultivation of viruses
CO3	Able to understand the cultivation of soil microbes
CO4	Able to understand the water quality parameter techniques
CO5	To understand the isolation of bacteria from spoiled fruits

- 1. Haemagglutination.
- 3. Egg inoculation methods (Demostration).
- 4. Wet mount examination of parasites.
- 5. Concentration methods for egg / ova
  - Flotation technique
  - Sedimentation technique
- 6. Isolation of bacteria from rhizosphere.
- 7. Plant diseases Fungi and Bacteria.
- 8. MPN and Settle Plate method.
- 9. 10. Dissolved oxygen.
- 11. MBRT and Resazurin test.
- 12. Isolation of bacteria from spoiled fruits and soft drinks.

# Mapping

CO/PSO	PSO1	PSO2	PSO3	PSO4	PSO5
CO1	✓	✓	✓	✓	✓
CO2	✓	✓	✓	✓	✓
CO3	✓	✓	✓	✓	✓
CO4	✓	✓	✓	✓	✓
CO5	✓	✓	✓	✓	✓