



## Integral University, Lucknow

<b>Effective from Session: 2023-24</b>							
<b>Course Code</b>	B100301T/ BS207	<b>Title of the Course</b>	Molecular Biology	<b>L</b>	<b>T</b>	<b>P</b>	<b>C</b>
<b>Year</b>	II	<b>Semester</b>	III	<b>3</b>	<b>1</b>	<b>0</b>	<b>4</b>
<b>Pre-Requisite</b>	10+2 Biology	<b>Co-requisite</b>					
<b>Course Objectives</b>	The objective of this course is to enable students to understand the concept of different types of genes, DNA replication, Transcription, Translation, regulation of Gene expression in prokaryotes and eukaryotes.						

### Course Outcomes

<b>CO1</b>	The students will be able to explain the basic concept of genome organization.
<b>CO2</b>	The students will be able to explain the process of DNA replication and its regulation in prokaryotes and eukaryotes
<b>CO3</b>	The students will be able to explain the process of transcription in prokaryotes and eukaryotes and post transcriptional modifications
<b>CO4</b>	The students will be able to describe the basics of translation in prokaryotes and eukaryotes and post translational modification
<b>CO5</b>	The students will be able to discuss regulation in gene expression and DNA repair systems.

Unit No.	Title of the Unit	Content of Unit	Contact Hrs.	Mapped CO
1	Basic Concepts of genome and its organization	Importance of Molecular Biology, Nucleic acid as the genetic material, Central Dogma of Molecular Biology, Model organisms for studying Molecular Biology, Genome and its organization in prokaryotes and Eukaryotes: Gene, Genome, Exon, Intron, regulatory sequence, Nucleosome structure and packaging of DNA into higher order structures.	8	CO1
2	DNA Replication	Semiconservative mode of replication. Mechanism of Replication in prokaryotes and eukaryotes. Enzymes and proteins involved in replication, Theta model and Rolling circle model, Inhibitors of Replication.	8	CO2
3	DNA Damage, Repair and Mutation	Causes and types of DNA damage, Mechanism of DNA repair, Molecular basis and types of mutation. Ames test.	6	CO3
4	Transcription	Transcription process in prokaryotes and eukaryotes. Enzymes, promoter, and transcription factors. Inhibitors of transcription Actinomycin D and $\alpha$ - Amanitin.	8	CO4
5	Translation	Components of Protein synthesis machinery: Messenger RNA, tRNA structure and function, Charging of tRNA, aminoacyl tRNA synthetases, ribosome structure and assembly, Mechanism of protein synthesis in prokaryotes and Eukaryotes.	8	CO4
6	Post-Transcription and Post-Translation Modifications	Post-transcriptional modifications of eukaryotic mRNA (capping, polyadenylation and splicing, post-translational modifications of proteins.	8	CO5
7	Gene expression	Principles of gene regulation, negative and positive regulation, concept of operons, Regulation of gene expression in prokaryotes and eukaryotes; Lac operon and Trp operon concept	8	CO5
8	Classes of DNA sequences	Satellite DNA, Split genes, Pseudogenes, Transposable elements, Retroelements, LINES, SINEs.	6	CO5

#### Reference Books:

1. Lewin B. (2000). Genes VII. Oxford University press.
2. Watson JD, Hopkins NH, Roberts JW, Steitz JA, Weiner AM. (1987). Molecular biology of the gene.
3. Lodish H, Baltimore D, Berk A, Zipursky SL, Darnell J. (1995). Molecular cell biology.
4. Brown, TA Genomes (2020).
1. Lewin B. (2000). Genes VII. Oxford University press.

#### e-Learning Source:

PO-PSO CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PSO1	PSO2	PSO3	PSO4
<b>CO1</b>	3	1					1	2	2	1	
<b>CO2</b>	3	1					1	3	2	2	
<b>CO3</b>	3	1					1	3	2	3	
<b>CO4</b>	3	1					1	3	2	3	
<b>CO5</b>	3	1					1	3	2	3	

**1- Low Correlation; 2- Moderate Correlation; 3- Substantial Correlation**

<b>Name &amp; Sign of Program Coordinator</b>	<b>Sign &amp; Seal of HoD</b>
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## Integral University, Lucknow

<b>Effective from Session: 2023-24</b>							
<b>Course Code</b>	B100303T/ BS208	<b>Title of the Course</b>	Basics of Microbiology	<b>L</b>	<b>T</b>	<b>P</b>	<b>C</b>
<b>Year</b>	II	<b>Semester</b>	III	<b>3</b>	<b>1</b>	<b>0</b>	<b>4</b>
<b>Pre-Requisite</b>	10+2	<b>Co-requisite</b>					
<b>Course Objectives</b>	<b>The objective of this course is to develop an understanding of basics of microbiology and sterilization techniques</b>						

### Course Outcomes

<b>CO1</b>	To understand basic details of basics and history of microbiology and general classification of microbes and extremophiles
<b>CO2</b>	To understand basic details of microbes in extreme environments and microbial interactions
<b>CO3</b>	To understand basic details of control of Microorganisms and staining techniques
<b>CO4</b>	To understand basic details of growth of microbes and recombination in Prokaryotes
<b>CO5</b>	To understand basic details of bacteriophages

Unit No.	Title of the Unit	Content of Unit	Contact Hrs.	Mapped CO
1	History of microbiology	Definition and scope of microbiology, Importance of microbiology in various fields, History of microbiology: Spontaneous generation and its controversy, Louis Pasteur and the refutation of spontaneous generation, Germ theory of disease, Robert Koch and the postulates of bacterial pathogenesis	8	CO1
2	Classification of microbes	Introduction to Microbial Classification, Prokaryotic, and eukaryotic microbes, Classification Methods and Techniques: Phenotypic, genotypic and serological methods, Microbial Taxonomy and Nomenclature; Nature of the microbial cell surface, gram positive and gram negative bacteria	8	CO1
3	Microbes in extreme environments and microbial interactions	Microbes in extreme environments and microbial interactions: The thermophiles: alkalophiles, acidophiles and symbiosis and antibiosis among microbial population, N <sub>2</sub> fixing microbes in agriculture and forestry.	8	CO2
4	Control of Microorganisms	Control of Microorganisms: Physical agents (Autoclave, Hot air oven, Laminar airflow and membrane filter.), chemical agents (Alcohol, Halogens and Gaseous agents, antibiotics), Radiation Methods (UV rays).	8	CO3
5	Stains and staining techniques	Introduction to Stains and Staining Techniques, Principles of staining, Types of stains – simple stains, structural stains, and Differential stains, Application of Staining Techniques in Microbial Diagnostics	6	CO3
6	Recombination in Prokaryotes	Recombination in Prokaryotes: Transformation, Conjugation and Transduction	8	CO4
7	Growth of microbes	Introduction to Microbial Growth, Microbial Growth Curve, Factors Influencing Microbial Growth	6	CO4
8	Viruses/Bacteriophage	Introduction to Bacteriophages, Bacteriophage Structure and genetics, Bacteriophage Life Cycle: Lytic and lysogenic cycle, General characteristics of plant and animal viruses	8	CO5

#### Reference Books:

1. Introduction to Microbiology, Ingraham, 2ed.
2. Brock Biology of Microorganisms, Madigan et al, 9th ed.
3. General Microbiology, R.Y. Stanier, J.L. Ingraham, M.L.Wheelis and P.R. Painter, Macmillian
4. Microbiology VI Edition, M.J. Pelczar, E.C.S. Chan and N.R. Kreig, Tata McGraw Hill
5. Principles of Microbiology, R.M. Atlas, Wm C. Brown Publisher.
6. The Microbial World, Roger Y. Stanier, Prentice Hall
7. Howe.C. (1995) Gene Cloning and manipulation, Cambridge University Press, USA
8. Lewin, B., Gene VI New York, Oxford University Press.

#### e-Learning Source:

<https://www.khanacademy.org/>

### Course Articulation Matrix: (Mapping of COs with POs and PSOs)

PO- PSO CO	Course Articulation Matrix: (Mapping of COs with POs and PSOs)											
	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PSO1	PSO2	PSO3	PSO4	
CO1	3	1					1	3		1	2	
CO2	3	1					1	3		2	2	
CO3	3	1					1	3		3	2	
CO4	3	1					1	3		3	2	
CO5	3	1					1	3		3	2	

1- Low Correlation; 2- Moderate Correlation; 3- Substantial Correlation

Name & Sign of Program Coordinator	Sign & Seal of HOD
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## Integral University, Lucknow

<b>Effective from Session: 2023-24</b>							
<b>Course Code</b>	B190302P /BS209	<b>Title of the Course</b>	Molecular Biology Lab	<b>L</b>	<b>T</b>	<b>P</b>	<b>C</b>
<b>Year</b>	II	<b>Semester</b>	III	<b>0</b>	<b>0</b>	<b>4</b>	<b>2</b>
<b>Pre-Requisite</b>	10+2	<b>Co-requisite</b>					
<b>Course Objectives</b>	The course is designed to train the students in basic and some advanced techniques of Molecular biology.						

Course Outcomes	
<b>CO1</b>	The students will be able to isolate genomic DNA from bacteria.
<b>CO2</b>	The students will be able to isolate genomic DNA from plant or animal tissues.
<b>CO3</b>	The students will be able to isolate plasmid DNA ( <i>E. coli</i> ).
<b>CO4</b>	The students will be able to perform restriction digestion of DNA.
<b>CO5</b>	The students will be able to perform Agarose Gel Electrophoresis.

Unit No.	Title of the Unit	Content of Unit	Contact Hrs.	Mapped CO
1	Exp-01	Isolation of genomic DNA from bacteria ( <i>E. coli</i> )	4	CO1
2	Exp-02	Isolation of genomic DNA from plant tissue	6	CO2
3	Exp-03	Isolation of genomic DNA from animal tissue	6	CO2
4	Exp-04	Isolation of plasmid DNA ( <i>E. coli</i> )	4	CO3
5	Exp-05	Restriction digestion of DNA	2	CO4
6	Exp-06	Agarose Gel Electrophoresis	2	CO5

**Reference Books:**

1. Gene Cloning and DNA Analysis: An Introduction, 6th Edition by T. A. Brown
2. Sambrook J, Russell D (2001) Molecular Cloning: A Laboratory Manual, 3rd Ed. Cold Spring Harbor, NY: Cold Spring Harbor Laboratory Press.

**e-Learning Source:**

**Course Articulation Matrix: (Mapping of COs with POs and PSOs)**

PO-PSO CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PSO1	PSO2	PSO3	PSO4
<b>CO1</b>	3	1						3		3	
<b>CO2</b>	3	1		3		3	1	3	2	3	
<b>CO3</b>	3	1		3		3	1	1		3	
<b>CO4</b>	3	1		3		3	1				3
<b>CO5</b>	3	1		3	3	3	1				3

1- Low Correlation; 2- Moderate Correlation; 3- Substantial Correlation

Name & Sign of Program Coordinator	Sign & Seal of HOD
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## Integral University, Lucknow

<b>Effective from Session: 2023-24</b>							
<b>Course Code</b>	B100304P/ BS210	<b>Title of the Course</b>	Microbiology Lab	<b>L</b>	<b>T</b>	<b>P</b>	<b>C</b>
<b>Year</b>	II	<b>Semester</b>	III	<b>0</b>	<b>0</b>	<b>4</b>	<b>2</b>
<b>Pre-Requisite</b>	10+2	<b>Co-requisite</b>					
<b>Course Objectives</b>	The objective of this course is to develop the understanding of basic microbiology, Instruments used to study and work on microbes, Staining Techniques, Enzyme assay and Biochemical tests–starch hydrolysis, gelatin liquefaction, Cleaning and sterilization of glassware, Media preparation and Isolation of bacteria and fungi from various sources, Growth curve of bacteria, Isolation and purification and estimation of DNA and RNA						

Course Outcomes	
<b>CO1</b>	Develop an understanding of Instruments: Compound microscope, Autoclave, Hot air oven, pH meter, Laminar airflow and centrifuge.
<b>CO2</b>	Develop an understanding staining techniques
<b>CO3</b>	Understand processes involved in culturing of microbes as cleaning and sterilization of glassware, media preparation.
<b>CO4</b>	Understand the process of isolation of bacteria and fungi from soil/ air/water/ other sources
<b>CO5</b>	Understand the growth pattern of bacteria.

Unit No.	Title of the Unit	Content of Unit	Contact Hrs.	Mapped CO
1	Exp 1	Study of instruments: Compound microscope, Autoclave, Hot air oven, pH meter, Laminar airflow and centrifuge	8	CO1
2	Exp 2	Cleaning and sterilization of glassware	4	CO3
3	Exp 3	Media preparation: Nutrients agar, Nutrient broth and LB.	4	CO3
4	Exp 4	Isolation of bacteria and fungi from soil/ air/water – dilution and pour plate methods	8	CO4
5	Exp 5	Staining Techniques: Gram staining for gram positive and gram negative bacteria	8	CO2
6	Exp 6	Growth curve of bacteria	8	CO5

**Reference Books:**

- Keith Wilson John Walker John M. Walker “Principles and Techniques of Practical Biochemistry”
- William M., Ph.D. O’Leary Robert Dony Wu “Practical Handbook of Microbiology”
- Joseph Sambrook David W. Russel Joe Sambrook “Molecular Cloning: A Laboratory Manual”

**e-Learning Source:**

**Course Articulation Matrix: (Mapping of COs with POs and PSOs)**

PO-PSO CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PSO1	PSO2	PSO3	PSO4
CO1	3	1		3		3	1	3	2	3	
CO2	3	1		3		3	1	1		3	
CO3	3	1		3		3	1				3
CO4	3	1		3	3	3	1				3
CO5	3	1		2		2	1				2

**1- Low Correlation; 2- Moderate Correlation; 3- Substantial Correlation**

Name & Sign of Program Coordinator	Sign & Seal of HOD
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## Integral University, Lucknow

**Effective from Session: 2023-24**

<b>Course Code</b>	B100303V/ BS247	<b>Title of the Course</b>	Molecular Diagnostics	<b>L</b>	<b>T</b>	<b>P</b>	<b>C</b>
<b>Year</b>	II	<b>Semester</b>	III	<b>3</b>	<b>0</b>	<b>0</b>	<b>3</b>
<b>Pre-Requisite</b>	10+2	<b>Co-requisite</b>					
<b>Course Objectives</b>	The objective of this course is to develop an understanding of the basic principle and application of molecular techniques employed in diagnosis of diseases.						

### Course Outcomes

<b>CO1</b>	To gain the basic knowledge about mechanism and pathogenesis of common diseases.
<b>CO2</b>	To understand basic details of pathogenesis and diagnosis of infectious diseases caused by bacteria, fungi, virus, and protozoa.
<b>CO3</b>	To understand basic details of basic principle & application of classical genotyping techniques.
<b>CO4</b>	To understand basic details of types of cancers, genetics and types of cancer and applications of Molecular Diagnostics for Human Cancers.
<b>CO5</b>	To understand basic details of principle and application of Molecular diagnostics techniques such as PCR, Real- Time PCR, DNA Sequencing, Microarray etc.

Unit No.	Title of the Unit	Content of Unit	Contact Hrs.	Mapped CO
1	Introduction to Human Genome & common diseases	Introduction and mechanism related to the human genome, such as gene expression, replication and genome maintenance. Consequences of mutations and polymorphisms, and impacts of genes and environment on major common diseases, such as cancer, diabetes, vascular and coronary disease	10	CO1
2	Infectious Diseases and History of Diagnostics	Types of infectious diseases- bacterial, viral, fungal, protozoans and other parasites. Infection mode of transmission in infections, factors predisposing to microbial pathogenicity. Diagnosis of infectious diseases caused by bacteria, fungi, viruses, protozoa and Helminthes.	10	CO2
3	Major Genetic disorders, its causes & Diagnosis.	Genetic disorders; Sickle cell anaemia, Duchene muscular Dystrophy, Retinoblastoma, Cystic Fibrosis and Sex – linked inherited disorders	10	CO3
4	Cancer Biology and Diagnostics	Different types of cancers, genetics of cancer- oncogenes, tumour suppressor genes, Applications of Molecular Diagnostics for Human Cancers.	8	CO4
5	Molecular Diagnostics Tools	RT- PCR, Animal cell culture, DNA Sequencing, Microarray, Techniques of Nucleic acid Extraction, Real time PCR, Fluorescence In Situ Hybridization.	7	CO5

**Reference Books:**

- "Murray's Medical Microbiology" by Patrick R. Murray, Ken S. Rosenthal, Michael A. Pfaller
- "Medical Microbiology" by David Greenwood, Richard C. B. Slack, Michael R. Barer, Will L. Irving
- "Kuby Immunology" by Judy Owen, Jenni Punt, Sharon Stranford
- "Basic Immunology: Functions and Disorders of the Immune System" by Abul K. Abbas, Andrew H. Lichtman

**e-Learning Source:**

Vlab.amrita.edu

### Course Articulation Matrix: (Mapping of COs with POs and PSOs)

PO- PSO CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PSO1	PSO2	PSO3	PSO4
<b>CO1</b>	3	1	1				1	3		1	3
<b>CO2</b>	3	1					1	2		2	3
<b>CO3</b>	3	1		3			1	3		3	2
<b>CO4</b>	3	1					1	2		3	2
<b>CO5</b>	3	1					1	3		3	2

1- Low Correlation; 2- Moderate Correlation; 3- Substantial Correlation

Name & Sign of Program Coordinator	Sign & Seal of HOD
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## Integral University, Lucknow

<b>Effective from Session: 2023-24</b>							
<b>Course Code</b>	B100401T/ BS218	<b>Title of the Course</b>	Industrial Biotech and Bioprocess Technology	<b>L</b>	<b>T</b>	<b>P</b>	<b>C</b>
<b>Year</b>	II	<b>Semester</b>	IV	<b>3</b>	<b>1</b>	<b>0</b>	<b>4</b>
<b>Pre-Requisite</b>	10+2 Biology	<b>Co-requisite</b>					
<b>Course Objectives</b>	After completion of the course, a student will be able to develop the understanding of industrial aspects of biotechnology, IPR and bioethics						
<b>Course Outcomes:</b> After this course students will be able to							
<b>CO1</b>	To understand the problems in isolation, strain improvement in industrial processes.						
<b>CO2</b>	To understand the growth of microorganisms in industrial processes.						
<b>CO3</b>	To understand design and types of fermenters and operation of fermenters.						
<b>CO4</b>	To understand the production process of alcohols, antibiotic and enzymes and other biologically active compounds by industrial microbiological fermentation						
<b>CO5</b>	To understand the regulation of bioethics and policies of IPR and entrepreneurship.						

Unit No.	Title of the Unit	Content of Unit	Contact Hrs.	Mapped CO
1	Introduction	Introduction of Industrial microbiology and Bioprocess technology. History-Introduction, scope and relation with other sciences. Screening for new metabolites: primary and secondary products. Maintenance of strains. Strain development through selection, mutations and recombination, and other recent methods	8	CO1
2	Fermentation technology	Fermentation media, Natural and synthetic media, Sterilization techniques: Heat, Radiation and Filtration method. Types of fermentation: solid state, submerged fermentation and continuous fermentation, Types of microbial culture and its growth kinetics– Batch, Fedbatch and Continuous culture.	8	CO2
3	Bioprocess technology	Design and working of a typical bioreactor, Process of Aeration, Agitation, and Temperature regulation, Immobilized enzymes and cell bioreactors. Downstream processing (DSP), Disintegration of cells, Separation, Extraction, Concentration and purification of products.	8	CO3
4	Production of alcohols, antibiotic and enzymes:	Brief account of the following products obtained by industrial microbiological fermentation: alcohols (Ethanol) and Alcoholic Beverage: (Beer), Organic acid: (citric and acetic). Amino acids: (Glutamic acid), Vitamin: (vitamin B12).	8	CO3
5	Production of biologically active compounds:	Production of antibiotics (penicillin) and enzymes (amylase, protease). Production of microbial food and single cell proteins	8	CO4
6	IPR	Introduction to Intellectual Property Rights (IPR)-World Intellectual properties, Indian Intellectual Properties. Patents, Copyrights, Designs, Trademarks, Geographical Indication. Infringement of IPR, Its protection and Remedies. Licensing and its types.	7	CO5
7	Issues related to IPR	Issues related to IPR protection of software and database; IPR protection of life forms; patenting biological products and biodiversity; Major changes in Indian patent system as post TRIPS effects	6	CO5
8	Bioethics and GMP	Introduction, necessity and limitation; Different paradigms of bioethics: National and International; Ethical conflicts in Biotechnology; Bioethics of genes, Legal implications in bioethics. Introduction to GMP	7	CO5

**Reference Books:**

1. Glazier AN and Nikaido H (2007). Microbial Biotechnology – Fundamental & Applied Microbiology – Second Edition. Cambridge University Press.
2. Casida LE (2019) Industrial Microbiology. Second Edition, New Age International Publisher.
3. Stanbury P F and Whitaker, A. (2010). Principles of Fermentation Technology. Oxford: Pergamon Press
4. Shuler M L and Kargi F. (2002). Bioprocess Engineering: Basic Concepts. Upper Saddle River, NJ: Prentice Hall.
5. Crueger W and Crueger A (2002) Cruegers Biotechnology: A Textbook of Industrial Microbiology. Third Edition, Panima Publishing Corp., New Delhi.
6. Blanch H W and Clark D S. (1997). Biochemical Engineering. New York: M.Dekker.
7. Bailey J E and Ollis D F. (1986). Biochemical Engineering Fundamentals. New York: McGraw-Hill.

**e-Learning Source:**

<https://ocw.mit.edu/courses/civil-and-environmental-engineering/1-34-waste-containmentand-remediation-technology-spring-2004/lecture-notes/>  
<https://ocw.mit.edu/courses/civil-and-environmental-engineering/1-018j-ecology-i-theearth-system-fall-2009/>  
[https://ocw.mit.edu/courses/civil-and-environmental-engineering/1-018j-ecology-i-theearth-system-fall-2009/lecture-notes/MIT1\\_018JF09\\_Lec07.pdf](https://ocw.mit.edu/courses/civil-and-environmental-engineering/1-018j-ecology-i-theearth-system-fall-2009/lecture-notes/MIT1_018JF09_Lec07.pdf)  
<https://ocw.mit.edu/courses/civil-and-environmental-engineering/1-89-environmentalmicrobiology-fall-2004/>

PO-PSO CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PSO1	PSO2	PSO3	PSO4
CO1	3	1					1	3	2	3	
CO2	3	1					1	3	2	2	
CO3	3	1					1	3	2	3	
CO4	3	1					1	3	2	3	
CO5	3	1					1	3	2	3	

**1- Low Correlation; 2- Moderate Correlation; 3- Substantial Correlation**

<b>Name &amp; Sign of Program Coordinator</b>	<b>Sign &amp; Seal of HOD</b>
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## Integral University, Lucknow

<b>Effective from Session: 2023-24</b>							
<b>Course Code</b>	B100402P/ BS219	<b>Title of the Course</b>	Industrial Biotechnology Lab	<b>L</b>	<b>T</b>	<b>P</b>	<b>C</b>
<b>Year</b>	II	<b>Semester</b>	IV	<b>0</b>	<b>0</b>	<b>4</b>	<b>2</b>
<b>Pre-Requisite</b>	10+2 Biology	<b>Co-requisite</b>					
<b>Course Objectives</b>	The objective of this course is to enable students learn about basics of industrial biotechnology and fermentation						

Course Outcomes	
<b>CO1</b>	Understand method of isolation of industrially important microorganisms.
<b>CO2</b>	Perform Algal or fungal culture
<b>CO3</b>	Perform estimation of citric acid from <i>Aspergillus</i> culture.
<b>CO4</b>	Perform estimation of lactic acid.
<b>CO5</b>	Understand the working of small scale fermenter

Unit No.	Title of the Unit	Content of Unit	Contact Hrs.	Mapped CO
1	<b>Exp. 1</b>	Isolation of industrially important microorganisms from soil.	8	CO1
2	<b>Exp. 2</b>	Algal or fungal culture (Yeast and <i>Aspergillus</i> )	8	CO2
3	<b>Exp. 3</b>	Estimation of citric acid from <i>Aspergillus</i> culture.	8	CO3
4	<b>Exp. 4</b>	Estimation of lactic acid.	8	CO4
5	<b>Exp. 5</b>	Demo of working of small scale fermenter	8	CO5

<b>Reference Books:</b>	
1.	Glazier AN and Nikaido H (2007). Microbial Biotechnology – Fundamental & Applied Microbiology – Second Edition. Cambridge University Press.
2.	Casida LE (2019) Industrial Microbiology. Second Edition, New Age International Publisher.
3.	Stanbury P F and Whitaker, A. (2010). Principles of Fermentation Technology. Oxford: Pergamon Press
4.	Crueger W and Crueger A (2002) Crueger's Biotechnology: A Textbook of Industrial Microbiology. Third Edition, Panima Publishing Corp., New Delhi.
5.	Blanch H W and Clark D S. (1997). Biochemical Engineering. New York: M. Dekker.
<b>e-Learning Source:</b>	

PO-PSO CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PSO1	PSO2	PSO3	PSO4
	<b>CO1</b>	3	1		3			3	3	2	
<b>CO2</b>	3	1		3			3	3	2		3
<b>CO3</b>	3	1		3			3	3	2		3
<b>CO4</b>	3	1					3	3	2		3
<b>CO5</b>	3	1					3	3	2		3

**1- Low Correlation; 2- Moderate Correlation; 3- Substantial Correlation**

Name & Sign of Program Coordinator	Sign & Seal of HOD
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## Integral University, Lucknow

<b>Effective from Session: 2023-24</b>							
<b>Course Code</b>	B100403T/B S256	<b>Title of the Course</b>	Infection and immunity	<b>L</b>	<b>T</b>	<b>P</b>	<b>C</b>
<b>Year</b>	II year	<b>Semester</b>	IV sem	<b>3</b>	<b>1</b>	<b>0</b>	<b>4</b>
<b>Pre-Requisite</b>	10+2	<b>Co-requisite</b>					
<b>Course Objectives</b>	<b>The objective of this course is to develop an understanding of the basics of infection and immunity</b>						

Course Outcomes	
<b>CO1</b>	To understand basic details of infectious diseases and its transmission
<b>CO2</b>	To understand basic details of Laboratory Diagnosis of Infectious Agents
<b>CO3</b>	To understand basic details of antigens, immune system organization and types of immunity
<b>CO4</b>	To understand basic details of Structure and functions of MHC molecules and complement activation
<b>CO5</b>	To understand basic details of Immunological techniques, Immune response and Vaccination

Unit No.	Title of the Unit	Content of Unit	Contact Hrs.	Mapped CO
1	History and transmission of infectious diseases	Definition and Historical perspectives of infectious diseases, Modes of Transmission and Pathogenesis of Infectious Diseases: Adherence and invasion mechanisms, toxigenesis and virulence factors, Host-pathogen interactions	8	CO1
2	Laboratory Diagnosis of Infectious Agents	Laboratory Diagnosis of Infectious Agents: Sample collection and handling, Microscopic examination and staining techniques, Culture, biochemical tests, and serological assays; Infection Control Measures: Standard precautions and isolation techniques, Sterilization, disinfection, and decontamination, Surveillance and outbreak investigation	8	CO2
3	Immune system organization	History of Immune system, Types of immunity Humoral & Cell Mediated. The cells and organs of the immune system. Innate immunity. Anatomical barriers, cell types of innate immunity, connection between innate and adaptive immunity	8	CO3
4	Types of Immunity and antigenic determinants	Adaptive immunity: Antigens and haptens. Structure and distribution of classes and substances of immunoglobulins (Ig), Ig fold, effector functions of antibody, antigenic determinants on Ig and Ig super family. Generation of antibody diversity	8	CO3
5	Structure and functions of MHC molecules	Structure and functions of MHC molecules (MHC I and II), Endogenous and exogenous pathways of antigen processing and presentation	6	CO4
6	Complement and its activation	Complement and its activation by classical, alternate and lectin pathway; biological consequences of complement activation; regulation of complement activity	6	CO4
7	Immunological techniques	Immunological methods-Antigen-antibody interactions. Agglutination, hemagglutination. Precipitin reactions in solution and in gels; immunoassays. Selection, Antigen presentation, Activation of T and B cells. Cytokines	8	CO5
8	Immune response and Vaccination	Immunological tolerance-Primary and secondary. Hypersensitivity and its types. Immune response against major classes of pathogens. Vaccines: Live attenuated, Inactivated, Toxoid, subunit/conjugate vaccine. Monoclonal Antibody	8	CO5

**Reference Books:**

- "Murray's Medical Microbiology" by Patrick R. Murray, Ken S. Rosenthal, Michael A. Pfaller
- "Medical Microbiology" by David Greenwood, Richard C. B. Slack, Michael R. Barer, Will L. Irving
- "Kuby Immunology" by Judy Owen, Jenni Punt, Sharon Stranford
- "Basic Immunology: Functions and Disorders of the Immune System" by Abul K. Abbas, Andrew H. Lichtman

**e-Learning Source:**

Course Articulation Matrix: (Mapping of COs with POs and PSOs)											
PO-PSO CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PSO1	PSO2	PSO3	PSO4
CO1	3	1	1				1	3		1	3
CO2	3	1					1	2		2	3
CO3	3	1		3			1	3		3	2
CO4	3	1					1	2		3	2
CO5	3	1					1	3		3	2

1- Low Correlation; 2- Moderate Correlation; 3- Substantial Correlation

<p><b>Name &amp; Sign of Program Coordinator</b></p>	<p><b>Sign &amp; Seal of HOD</b></p>
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## Integral University, Lucknow

<b>Effective from Session: 2023-24</b>							
<b>Course Code</b>	B100404P/ BS258	<b>Title of the Course</b>	Immunological Techniques Lab	<b>L</b>	<b>T</b>	<b>P</b>	<b>C</b>
<b>Year</b>	II	<b>Semester</b>	IV	<b>0</b>	<b>0</b>	<b>4</b>	<b>2</b>
<b>Pre-Requisite</b>	10+2 Biology	<b>Co-requisite</b>					
<b>Course Objectives</b>	The objective of this course is to enable students learn about basics of immunology, types of Blood grouping, cell counts, ELISA, Ouchterlony Double diffusion (ODD) and Separation of serum from blood & precipitation of Immunoglobulins						

Course Outcomes	
<b>CO1</b>	Analyze Blood grouping
<b>CO2</b>	Perform and analyze differential counting of WBC and detergent lysis of RBC
<b>CO3</b>	Perform and analyze Dot Elisa, ELISA
<b>CO4</b>	Have knowledge of and can perform Ouchterlony Double diffusion assay
<b>CO5</b>	Perform and analyze separation of serum from blood & precipitation of Immunoglobulin

Unit No.	Title of the Unit	Content of Unit	Contact Hrs.	Mapped CO
1	<b>Exp. 1</b>	Blood grouping	6	CO1
2	<b>Exp. 2</b>	Differential Count of WBC	6	CO1
3	<b>Exp. 3</b>	Detergent lysis of RBC	6	CO2
4	<b>Exp. 4</b>	Dot Elisa	6	CO3
5	<b>Exp. 5</b>	ELISA – Demonstration	6	CO3
6	<b>Exp. 6</b>	Ouchterlony Double diffusion (ODD)	6	CO4
7	<b>Exp. 7</b>	Separation of serum from blood & precipitation of Immunoglobulins	6	CO5

**Reference Books:**

1. Asim Roy Kumar, 2. Talwar Gupta A Handbook of Practical & Clinical Immunology 3. A.K. Abbas and A.H. Lichtman, Saunders, Basic Immunology, W.B. Company

**e-Learning Source:**

PO-PSO CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PSO1	PSO2	PSO3	PSO4
CO1	3	1		3			2	3	2		
CO2	3	1		3			2	3	2		
CO3	3	1		3			2	3	2		
CO4	3	1					2	3	2		
CO5	3	1					2	3	2		

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<b>Name &amp; Sign of Program Coordinator</b>	<b>Sign &amp; Seal of HOD</b>
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## Integral University, Lucknow

**Effective from Session: 2023-24**

<b>Course Code</b>	B110405V/ BS259	<b>Title of the Course</b>	Molecular Medicine	<b>L</b>	<b>T</b>	<b>P</b>	<b>C</b>
<b>Year</b>	II	<b>Semester</b>	IV	<b>3</b>	<b>0</b>	<b>0</b>	<b>3</b>
<b>Pre-Requisite</b>	10+2	<b>Co-requisite</b>					
<b>Course Objectives</b>	The objective of this course is to develop an understanding of principle and application of the molecular medicine.						

### Course Outcomes

<b>CO1</b>	To understand basic knowledge of working, design, and requirements a molecular medicine lab set up along with sample handling and preparation in lab.
<b>CO2</b>	To understand basic understanding of conformations of Biomolecules and diseases related to protein mis- folding.
<b>CO3</b>	To understand basic details the principle and methodology employed for the studying tissue and cell structure, and different preparative procedures for light and electron microscopic visualization
<b>CO4</b>	To understand basic details about the principle and technical aspects of animal cell culture.
<b>CO5</b>	To understand basic details about principle and application of several molecular techniques employed in diagnosis of diseases.

Unit No.	Title of the Unit	Content of Unit	Contact Hrs.	Mapped CO
1	Introduction to Molecular Medicine Lab	Molecular Laboratory Set up: Introduction, Design, Requirements, Laboratory, Good Clinical Laboratory Practice (GCLP), buffer preparation, micro-pipetting, Measurement of pH of solutions, molarity, normality and molality calculation and graph plot, sample collection, handling and storage etc. used in laboratory.	8	CO1
2	Biomolecule Conformations & related disorders	Conformation of Biomolecules: Nucleic acids: A-, B-, Z-DNA forms. Ramachandran plot, Secondary, Tertiary and Quaternary structure, Domains, Motif and Folds. Protein misfolding: diseases and diagnosis	8	CO2
3	Cell Imaging and Interpretation	Principles and constituents of compound, fluorescence, phase contrast, differential interference contrast and dark field microscopy, Preparation of cells and tissues for light and electron microscopy.	8	CO3
4	Animal Cell Culture	Description and maintenance of animal cell culture, aseptic technique, cloning and selection of specific cell types, contamination, methods for measuring viability and cytotoxicity, cell culture environment (substrate, gas phase, medium) and the culturing of specific cell types	8	CO4
5	Molecular Diagnostics Techniques	Role of PCR & its variants in diseases diagnosis, Nucleic acid Extraction Protocol (DNA & RNA), Polymorphism based disease diagnostics techniques such as RFLP and RAPD.	6	CO5

**Reference Books:**

- " Berg, J.M., Tymoczko, J.L. and Stryer, L. (2010). Biochemistry. W.H. Freeman & Company. USA.
- "Medical Microbiology" by David Greenwood, Richard C. B. Slack, Michael R. Barer, Will L. Irving
- "Kuby Immunology" by Judy Owen, Jenni Punt, Sharon Stranford
- "Basic Immunology: Functions and Disorders of the Immune System" by Abul K. Abbas, Andrew H. Lichtman

**e-Learning Source:**

### Course Articulation Matrix: (Mapping of COs with POs and PSOs)

PO- PSO CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PSO1	PSO2	PSO3	PSO4
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<b>CO2</b>	3	1					1	2		2	3
<b>CO3</b>	3	1		3			1	3		3	2
<b>CO4</b>	3	1					1	2		3	2
<b>CO5</b>	3	1					1	3		3	2

1- Low Correlation; 2- Moderate Correlation; 3- Substantial Correlation

Name & Sign of Program Coordinator	Sign & Seal of HOD
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