										S	eme	ster –	· I								
Course code	2:								A	Allie	ed I	A				L	T	P	(7	H/W
23BMCA1						Bo	ody l	Flu	uid A	Ana	lysis	s					T		3		3
Objectives	> K > N	 Acquire knowledge of body fluids and their functions. Know about the Infection transmission process & its prevention Make aware of standard norms, principles, classification, sources & hazards associated with biomedical waste management. 																			
Unit –I	prope		body	flui																	ds. Physica cormalities of
Unit-II	Testin	Amniotic fluid: Formation and function of amniotic fluid, Chemical composition, Collection, Testing – Alpha fetoprotein, Acetyl cholinesterase, Neural tube defects, Chromosomal abnormalities, Haemolytic disease of newborn, Gestation age, Fetal maturation.																			
Unit III	exami	Cerebrospinal fluid:- Formation, Specimen collection, Causes of CSF pressure changes, Gross examination, Chemical analysis, Microbiologic examination, Immunologic tests, Cytological examination and clinical correlation and other fluid such as Serous fluid, Synovial fluid.																			
Unit IV	coagul & MC	Components of the blood (Plasma and Cellular elements) and their functions, Mechanism of coagulation of blood, Coagulation system, Haemogram, Calculations of Anaemia using MCH, MCV & MCHC, Special Haematological tests: Osmotic fragility – Heinz body preparation, Blood parasites – Lupus Erythematosus (LE)																			
Unit V	perfor		ate co	omple	lexit	ty T	Tests	ts: I	Princ	cipl	le, re	eportii									boratory the perform hig
Reference a Kanai Mukh			edical	l Lab	orat	atory	у Те	echr	nolo	gy,	volı	ıme –	I, II,	, III, T	`ataN	ЛсGı	aw F	Iill.			
Praful.B. Go Publ	dkar, e ication	tal., (1996 Hoxe	6). Te	extbo	ook (of N	Med	edica	al La	aboı	rator	у Тес	hnol	ogy, 2	2 nd ed	lition	, Bha	ılani			
Sambrook J Sprii		ssell DW, oratory Pre					ılar o	clor	ning	g — <i>F</i>	A lab	orato	ry m	anual	, 3 rd	editio	on, V	ol. I -	- III,	, Co	old
Outcomes After completion of the course, student Know the routes of infectious a Acquire knowledge on sterilizat Manage the biomedical waste.						s age zatio	ents'	' trans	miss	sion a			con	rol tl	ne di	sea	ses.				

		Semester - I							
Course code:		Practical I A	T	P	C	H/W			
23BMCAP1	Ī	Lab in Body Fluid Analysis			P	2	2		
Objectives	A A A	Determine the levels of body fluids and know their functions. Know about the Infection transmission process & its prevention Make aware of standard norms, principles, classification, sources & hazards associated with biomedical waste management.							
1. Standardization of distilled or deionized water.									
2. Microscopic examination of total leukocyte count.									

- 3. Determination of serum alkaline phosphatase by PNP method.
- 4. Determination of urine creatinine
- 5. Perform serological diagnosis of microbial diseases
- 6. Anti-streptolysin O (ASO) quantitative test
- 7. Perform C- reactive protein test (CRP)
- 8. Determination of blood hemoglobin by cyanomethemoglobin method
- 9. Reference ranges and normal values of RBC, Haemoglobin, WBC, Differential white cell count.
- 10. Hemorrhagic disorders related to platelet and capillary defects.

Grimaldi and Scopacasa (2000) 'Evaluation of the Abbott CELL-DYN 4000 Hematology Analyzer', American Journal of Clinical Pathology.

Kanai Mukherjee, (2000). Medical Laboratory Technology, volume – I, II, III, Tata McGraw Hill.

Technology, 2nd Praful.B. Godkar,et (1996). Textbook Medical Laboratory al., of edition, Bhalani Publication House

After completion of the course, students are expected to be able to: **Outcomes**

- > Determine the leukocyte count, urea creatinine and blood hemoglobin.
- > Identification of antigens by serological tests.
- Acquire basic knowledge on the reference and normal values of RBC and WBC.

		Semester - II					
Course coo	le: 23BMCA2	Allied I B	L	T	P	C	H/W
		Blood Banking Technology		Т		3	3
Objectives	The impression						
Unit –I Unit-II	Blood donation: Donor Motivation, Motivational Techniques, Social Marketing, Preparation of IE Materials. Donor recruitment & Retention: Types of blood donors, Donor selection, medical intervie and medical examination, screening for haemoglobin estimation, Managingrejected blood donors, technique for conversion of first time donor into regular voluntary donor, donor felicitation. Blood collection room equipment, their principles, and use, emergency medicines, Pre donation counselling, Bleeding of the donor, post donation care, post donation counseling. Screening of blood units for mandatory test Discarding infected units. Blood Banking- Blood Components: Selection of blood bags for component preparation, preparation of recell concentrate, Fresh Frozen plasma, platelet concentrate, cryoprecipitate, washed red cells, Frozen recells. Plasma Fractionation: Principles, manufacturing of different plasma derivatives- Component Testing Labeling - Transportation and storage of blood components.						
Unit III							
Unit IV	Quality Control Documentation and Legal Aspects of Blood Banking: Quality control of blood grouping reagents, QC of anti-human globulin reagent, bovine albumin, Normal saline- Quality control of blood bags -Quality control of different blood bank Components, sterility test on component - Organization of blood bank services, Blood Bank premises and infrastructure, Regional blood transfusion centre and blood storage centres.						
Unit V	Recent Advances In - Apheresis - Stem Ce	Blood Banking Techniques: Automation in lls.	Blood 1	Banki	ing -	Nuclei	ic Acid Testing
D 6	170 (1 1	·					

Abbas A K and Lichtman. A H. Basic Immunology, Saunders Elsevier.

David Latchman, 1997.Basic molecular and cell biology.BMJ Publishing group.

Denise M Harmening. Modern Blood Banking and Transfusion practices by, (5th ed)

Mollison PL Dacie, J A and Lewis S M Blood transfusion in clinical medicine-Practical Hematology.

National guide book in blood donor motivation. NACO, Ministry of Health and Family Welfare, Govt. of India.

Roitt, I. Essential Immunology. (8th ed), Blackwell scientific publications

Standards for blood banks and blood transfusion services, NACO, 2007. Ministry of Health and Family Welfare, Govt. of India, New Delhi.

Transfusion Medicine technical manual-DGHS, 2003. Ministry of Health and Family Welfare, Govt. of India (2nd ed)

Voluntary blood donation program NACO, 2007. Ministry of Health and FamilyWelfare, Govt. of India, New Delhi,

Outcomes After completion of the course, students are expected to be able to: ➤ Acquire depth knowledge of selecting suitable blood donor and analysis of the blood components. ➤ Know how to maintain the blood collection bags and preparation of blood for transfusion. ➤ Be able to access the recent advance in blood banking techniques.

	Semester - II						
Course code: 23BMCAP2		Practical I B	L	T	P	C	H/W
		Lab in Blood Banking Technology			P	2	2
Objectives	➤ The im	wledge on of blood banking. pression of the transfusion therapy. cent advances in the blood banking techniques.					

- 1. Qualitative test for ABO grouping with antisera and tube method
- 2. Cross reactivity
- 3. Coomb's test-direct and indirect method
- 4. Confirmation of HIV 1 and 2 using ELISA
- 5. VDRL test for the confirmation of syphilis
- 6. Examination of Plasmodium in blood by leishman staining
- 7. Isolation of DNA from blood
- 8. Demonstration for the confirmation of Hepatitis B and C

David Latchman, 1997. Basic molecular and cell biology. BMJ Publishing group.

Mollison PL Dacie, J A and Lewis S M Blood transfusion in clinical medicine- Practical Hematology.

National guide book in blood donor motivation. NACO, Ministry of Health and Family Welfare, Govt. of India.

Standards for blood banks and blood transfusion services, NACO, 2007. Ministry of Health and Family Welfare, Govt. of India, New Delhi.

Transfusion Medicine technical manual-DGHS, 2003. Ministry of Health and Family Welfare, Govt. of India (2nd ed)

Voluntary blood donation program NACO, 2007. Ministry of Health and Family Welfare, Govt. of India, New Delhi,

Outcomes	After completion of the course, students are expected to be able to:
	Acquire depth knowledge of selecting suitable blood donor and analysis of the
	blood components.
	➤ Know how to maintain the blood collection bags and preparation of blood for transfusion.
	➤ Be able to access the recent advance in blood banking techniques.

Course code:	23PMC A 2	Semester - III Allied II A	L	Т	P	C	H/W				
Course coue:	ZSDNICAS	Hospital infection Control Practices	L	T	Г	3	3				
01: "	> TT 1 4	_	<u>,</u>	_	1.						
Objectives	> Understand the healthcare-associated infections & infection control policies										
	> Know about the Infection transmission process & its prevention										
		Make aware of standard norms, principles, classification, sources &hazards									
Unit –I		ed with biomedical waste management.	ation a	ontu	al my	0.000	n. Introduction				
Umt –I		Introduction of healthcare-associated infections & infection control program: Introduction,									
		Role & responsibilities of ICN, Role of hospital administration in hospital infection control,									
	Infection Protection for Healthcare Workers, Education and training of healthcare workers, patients, and families.										
Unit-II			& vari	0118	route	es of	transmission of				
	Infection transmission & its prevention: Introduction & various routes of transmission of infection, Standard / Universal precautions and its components, The significance of taking										
	standard/ Universal precautions, Isolation policies and procedures and Infection Control										
		measures to Control Transmission.									
Unit III	Sterilization and disinfection: Physical and chemical methods of sterilization and disinfection										
	Cleaning & Disinfection of medical equipment, Disinfection of Hepatitis B virus, Hepatitis C										
	virus, HIV or TB contaminated devices.										
Unit IV	Personal protective equipment and standard precautions: Introduction, Types & Method of										
	use of personal protective equipment (PPE): Gloves, Gown, mask, apron Protective eyewea										
	(goggles), Boots or shoe cover & Cap or hair cover. Hand hygiene practices: Introduction, type										
	of hand washing, Steps of hand washing, The role of hand hygiene in control of hospital-acquired.										
Unit V		waste management: Introduction, Standard n					· .				
		lanagement, WHO Classification of BMWN	*								
		ociated with biomedical waste management,		ls re	lated	to bi	omedical waste				
		, Treatment and disposal techniques of BMWM	1.								
Reference an			1.1 1			*******					
		- Prevention and Control by Purva Mathur. Pub	olisher:L	_ippi	ncott	W III1	ams				
&Will											
National, CD	C, WHO guidel	lines on Hospital Infection Control.									
Journals:											
\ T	1 CTT : 17	0 1									

- Journal of Hospital Infection.
 Journal of patient safety and infection control.
 American Journal of Infection Control.
 Waste Management Journal Elsevier.

L	/ Waste Wat	Waste Wanagement Journal Lisevier.							
	Outcomes	After completion of the course, students are expected to be able to:							
		➤ Know the routes of infectious agents' transmission and how to control the diseases.							
		➤ Acquire knowledge on sterilization and disinfection.							
		➤ Manage the biomedical waste.							

		Semester – III					
Course code:		Practical II A	L	T	P	C	H/W
23BMCAP3		Lab in Hospital Infection Control Measures			P	2	2
Objectives >		Know the basic techniques followed in the hospital & diseases Acquire knowledge in the identification of infection measures Perform basic and serological tests for the disease of	us aį	gents	and la		

- 1. Organization of infection control and surveillance of hospital acquired infections.
- 2. Precaution measures for nosocomial infections
- 3. Examination of Hand Hygiene
- 4. Laboratory first aid measures
- 5. Preparation of normal saline
- 6. Examination of decontamination of Hospital Environment
- 7. Prevention of Device Associated Infections
- 8. Preventive Strategies for Surgical Site Infections
- 9. Examination of morphology of blood cells
- 10. Determination of bleeding time
- 11. Determination of blood clotting time by capillary method and Lee- White method
- 12. Antibiotic sensitivity test by disc diffusion method
- 13. Various culture media used for mycotic organisms

- 1. Anudita Bhargava, Atul Jindal, etc. (2019). Hospital infection Control Measures, All India Sciences of Medical Institute, Raipur.
- 2. Hospital Infection Control Manual, (2017). Sigma Hospital, India.
- 3. Praful.B. Godkar, et al., (1996). Textbook of Medical Laboratory Technology, 2nd edition, Bhalani Publication House
- 4. Kanai Mukherjee, (2000). Medical Laboratory Technology, volume I, II, III, Tata McGraw Hill

Outcomes After completion of the course, students are expected to be able to: ➤ Do the first aid ➤ Know how to prevent the environment and patients in the hospital from infections by applying various techniques learned through this course. ➤ Acquire knowledge on basic tests followed in the hospital such as calculation of bleeding time and clotting time.

		Semester - IV						
Course code:	23BMCA4	Allied II B		L	T	P	С	H/W
		Microbial Biotechnol	ogy		T		3	3
Objectives	 Provide the student with an understanding of the current views of microbial association in various environments. Evaluate the continuing roles played by microbes in the environment. Recognition of microorganisms as indicators of alteration of an ecosystem. Understand microbial processes aimed to solve environmental problems. 							
Unit –I	Brief history of fermentation; Fermentation- general concepts, Applications of fermentation; Range of fermentation process- Microbial biomass, enzymes, metabolites, recombinant products, transformation process; Component parts of a fermentation process.							
Unit-II	Microbial biotechnology: Scope and its applications in human therapeutics, agriculture (Biofertilizers, PGPR, Mycorrhizae), environmental, and food technology, Use of prokaryotic and eukaryotic microorganisms in biotechnological applications, Genetically engineered microbes for industrial applications: Bacteria and yeast							
Unit III	acid; Gluconi	Organic feedstock: ethanol; Acetone; Ethanol Organic acids: Production of Citric acid; Acetic acid; Lactic acid; Gluconic acid; Kojic acid; itaconic acid; Amino acids: Use of amino acids in industry; methods of production; Production of individual aminoacids (L-Glutamic acid; L Lysin; L-Tryptophan).						
Unit IV	Enzymes: commercial applications; production of Amylases; Glucose Isomerase; L Asparaginase Proteases Renin; Penicillin acylases; Lactases; Pectinases; Lipases; Structure and biosynthesis Nucleosides Nucleotides and related compounds.							
Unit V	Vitamins- Vitamin B12; Riboflavin; B carotene; Antibiotics: beta-Lactam antibiotics; amino acid and peptide antibiotics; Carbohydrate antibiotics; Tetracycline and antracyclines; Nucleoside antibiotics; Aromatic antibiotics; bioplastics (PHB; PHA); biotransformation of steroids.							

Crueger Wand Crueger, A. Biotechnology. A Textbook of Industrial Microbiology, Sinauer Associates Publisher

Reed, G. *Industrial microbiology*, CBS publications

Demain L Biology of Industrial microorganisms, Stanbury P.F.A

Vogel H C, Todaro C.L, Todaro C.C. Fermentation and Biochemical Engineering Handbook: Principles, Process Design, and Equipment, Noyes Data Corporation/ Noyes Publications.

Scheper. T, New Products and New Areas of Bioprocess Engineering (Advances in Biochemical Engineering/Biotechnology, 68) Springer Verlag Publications

Outcomes	After completion of the course, students are expected to be able to:						
	Understand on soil characteristics and biogeochemical cycling						
	➤ Be familiar with the microbial analysis of drinking water and						
	Aeromicrobiology						
	➤ Know the different aspects of waste management and sewage Treatment systems						
	Acquire knowledge on bioremediation and microbial leaching						

	Semester - IV							
Course code: 23BMCAP4		Practical II B	L	T	P	C	H/W	
		Lab in Microbial Biotechnology			P	2	2	
Objectives	➤ Impart kr	 Highlight the roles and characteristics of microorganisms in field of Biotechnology Impart knowledge on the basic concept of multiplication in microorganism Know the metabolic pathways and products can be used in biotechnology. 						

- 1. Isolation of industrially important microorganism from different sources using specificsubstrates.
- 2. Design and Preparation of Media for Bioprocesses.
- 3. Growth curve of bacteria/Yeasts in batch culture and calculation of maximum specificgrowthrate.
- 4. To study the various methods of biomass measurement.
- 5. Production of ethanol from sucrose by yeast.
- 6. Determination of yield coefficient and Monod's constant and metabolic quotient of E.coli culture using glucose as a carbon source.
- 7. Design of fermenter.
- 8. Production of citric acid using sucrose and molasses.
- 9. Production of extracellular enzymes.
- 10. Ethanol production using immobilized yeast culture.

Atlas, R.M. and Bartha, R. 1992. *Microbial Ecology: Fundamentals and Applications*. (3rd ed) BenjaminCummings, Redwood City.CA.

Reed G, Industrial microbiology, CBS publications

Demain L Biology of Industrial microorganisms, Stanbury P.F.A

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Scheper. T, New Products and New Areas of Bioprocess Engineering (Advances in Biochemical Engineering/Biotechnology, 68) Springer Verlag Publications.

Outcomes	After completion of the course, students are expected to be able to:
	➤ Know the principles involved in preparation of Beverage and industrial Alcohols
	and the physical and chemical conditions influencing their production.
	➤ Understand the importance of microbial enzymes, their applications, production process and
	relate biotransformation principles to biotransformation of steroids
	➤ Conceptualize the principles and production process of different typesof Vaccines.