Savitribai Phule Pune University Syllabus for M. Sc. Microbiology Part II (Affiliated Colleges) (To be implemented from 2020-21) Course Structure, Semester III

Course Type	Course Code	Course Name	Credit	Assess	ment	
				IA	UA	Total
Core Compulsory Theory	CCTP 7	Immunology	4	30	70	100
Papers (CCTP)	(MB 701)					
	CCTP 8	Molecular Biology	4	30	70	100
	(MB 702)					
	CCTP 9	Clinical Microbiology	4	30	70	100
	(MB 703)					
Choice Based Optional	MBTE 31	Cell Culture Techniques	2	15	35	50
Papers (CBOP)	MBPE 31	Practicals based on Cell Culture	2	15	35	50
		Techniques				
Elective /Departmental		OR				
Course	MBTE 32	Bioremediation and Biomass utilization	2	15	35	50
	MBPE 32	Practicals based on Bioremediation and	2	15	35	50
		Biomass utilization				
	OR					
	MBTE 33	Microbial Virus Technology	2	15	35	50
	MBPE 33	Practicals based on Clinical	2	15	35	50
		Microbiology and Microbial Virus				
		Technology				
Core Compulsory	MBCP 3	Practicals based on Compulsory Theory	4	30	70	100
Practical Paper		Credits.				

Course Structure, Semester IV

Course Type	Course	Course Name	Credit	Assessment		
	Code			IA	UA	Total
Core Compulsory Theory Papers (CCTP)	CCTP 10 (MB 801)	Pharmaceutical Microbiology	4	30	70	100
	CCTP 11 (MB 802)	Microbial Technology	4	30	70	100
Any Two:	MBTE 41	Quality Assurance and Validation in Pharmaceutical Industry and Development of Anti-infectives	2	15	35	50
Choice Based Optional Papers (CBOP)	MBPE 41	Practicals based on quality assurance and validation in pharmaceutical industry and development of anti-infectives	2	15	35	50
Elective /Departmental				•		
Course	MBTE 42	Advances in Microbial Technology	2	15	35	50
	MBPE 42	Practicals based on Advances in Microbial Technology	2	15	35	50
	MBTE 43	Industrial Waste Water Treatment and Industrial Production of Vaccines	2	15	35	50
	MBPE 43	Practicals based on Industrial Waste Water Treatment and Industrial Production of Vaccines	2	15	35	50
	MBTE 44	Bioethics, Biosafety, Quality Control and Quality Assurance	2	15	35	50
	MBPE 44	Practicals based on Bioethics, Biosafety, Quality Control and Quality Assurance	2	15	35	50
Core Compulsory Practical Paper	MBCP 4	Dissertation	4	30	70	100

Savitribai Phule Pune University Syllabus restructuring 2020-2021 M.Sc. II Semester III

MB CCTP- 7 Immunology Core Compulsory Theory Paper

Total: 4 Credits

Workload:-15 hrs/credit

(Total Workload :- 4 credits x 15 hrs = 60 hrs in semester)

Credit	Credit Title and Contents	References
No.		
TC 1	Cell surface molecules and receptors	1. Austyn J. M. and Wood K. J. (1993) Principles of Molecular and Cellular Immunology.
	i. Definition, general Structure and mechanism	First edition Oxford University Press, New York.
	(dimerization and rotation), components of	2. Barret J. T. (1983) Text Book of Immunology. Fourth edition. Saint Louis, Mosby,
	signal transduction (extracellular signaling	London.
	molecule, receptor proteins, intracellular	3. Boyd W. C. (1966) Fundamentals of Immunology, Interscience Publishers, New York.
	signaling proteins and target proteins)	4. Gangal S. and Sontakke S. (2013) Textbook of Basic and Clinical Immunology.
	ii. Adhesion molecules in immune activation,	University Press, India.
	structure and function of B Cell Receptor,	5. Garcia K. C. and Adams E. J. (2005) How the T cell Receptor Sees Antigen - A
	TCR-CD3 complex, Toll-like receptors,	Structural View. Cell. 122(3): 333–336.
	Cytokine receptors, G-protein coupled	6. Hafler D. A. (2007) Cytokines and interventional immunology, Nature Reviews,
	receptors	Immunology. 7(6): 423-423.
	iii. Signal transduction pathways: IL-2 pathway	7. Kindt T. J., Osborne B. A. and Goldsby R. A. (2006) Kuby Immunology, Sixth edition,
	(JAK/STAT, Ras/MAP Kinase Pathways,	W. H. Freeman & Co.
	TCR-CD3 activation pathway)	8. Yoshimura A., Naka T. and Kubo M. (2007). SOCS proteins, cytokine signalling and
	1 27	immune regulation. Nature Reviews, Immunology, 7(6): 454-465.
TC 2	Regulation of Immune response	1. Abbas A. K. and Lichtman A. H. (2004) Basic Immunology. Functions and Disorders of
	i. Negative regulation - Immunological tolerance,	Immune System. Second edition. Elsevier Inc.
	Mechanisms of tolerance induction (related	2. Carroll M. C. (2004) The complement system in regulation of adaptive immunity. Nature
	experimentation using transgenic animals), T	Immunology. 5(10): 981-986.
	cell mediated suppression of immune response	3. Kindt T. J., Osborne B. A. and Goldsby R. A. (2006) Kuby Immunology. Sixth edition.
	ii. Regulation of immune responses by antigen,	W. H. Freeman & Co.

	antigen-antibody complexes, Network theory and its experimental evidence iii. Cytokine mediated cross regulation of TH subsets (TH1-TH2) iv. Regulation of complement system – Classical and alternative pathway v. Biological Response Modifiers for cancer therapy and autoimmune disorders	 Patwardhan B., Gautam M. and Diwanay S. (2006) Botanical Immunomodulators and Chemoprotectants in Cancer Therapy. In Drug discovery and development Volume I: Drug Discovery. Ed. Chorghade Mukund S. Wiley- Interscience, John Wiley and Sons Inc. USA. 405-424. Roitt I. M. (1984) Essentials of Immunology. P. G. Publishers Pvt. Ltd., New Delhi. Roitt I. M. 1988. Essentials of Immunology. ELBS, London. Yoshimura A., Naka T. and Kubo M. (2007) SOCS proteins, cytokine signalling and immune regulation. Nature Reviews. Immunology. 7(6):454-465
TC 3	 i. In vitro systems –Quantification of cytokines (ELISPOT assay), functional assays for phagocytes and cytokines (cytotoxicity and growth assays) ii. In vivo systems – Experimental animals in immunology research (Inbred animal strains, Knockout mice, transgenic animals), Animal models for autoimmunity and AIDS 	 Gangal S. and Sontakke S. (2013) Textbook of Basic and Clinical Immunology. University Press, India. House R. V. (1998) Therapeutic Manipulation of Cytokines, Biotechnology and Safety Assessment. Second edition. Taylor & Francis. 81-105. Kindt T. J., Osborne B. A. and Goldsby R. A. (2006) Kuby Immunology. Sixth edition. H. Freeman and Co. Mather J. P. and Roberts P. E. (1998). Introduction to Cell and Tissue Culture Theory and Technique. Plenum Publishing Corporation, New York. Roitt I., Brostoff J. and Male D. (1993) Immunology .Sixth edition .Mosby & Co. London. Talwar G. P. (1983). Handbook of Immunology. Vikas Publishing Pvt. Ltd. New Delhi. Paul W. E. (2003) Fundamental Immunology. 5th Ed. Lippincott. Williams and Wilkins Publishers.
TC 4	 Tumor Immunology i. Cellular transformations during neoplastic growth, Classification of tumors based on histological, Tumors of lymphoid system 	 Leen A. M., Rooney C. M. and Foster A. E. (2007) Improving T Cell Therapy for Cancer. Annu Rev. Immunol. 25 (1):243–65. Patwardhan B. Gautam M. and Diwanay S. (2006) Botanical Immunomodulators and Chemoprotectants in Cancer Therapy. In Drug discovery and development Volume I:
	(lymphoma, myeloma, Hodgkin's disease) ii. Escape mechanisms of tumor from host defence, Host immune response to tumor – Effecter mechanisms, Immuno- surveillance theory	 Drug Discovery. Ed. Chorghade Mukund S. Wiley- Interscience, John Wiley and Sons Inc. USA. 405-424. 3. Chatterjee C. C. (1992) Human Physiology Tenth edition Vol. 1 and 2. Medical Allied Agency, Calcutta. 4. Guyton A. C. and Hall J. E. (1996) Text Book of Medical Physiology. Goel Book

iii. Diagnosis of tumors – biochemical and immunological tumor markers iv. Approaches in cancer immunotherapy:	Agency, Bangalore. 5. Malati T. (2007) Tumor Markers: An Overview, Indian Journal of Clinical Biochemi 22(2):17-31.	istry.
Immune adjuvant and tumor vaccine ther		ation 00.

Molecular Biology II : CCTP-8 Core Compulsory Theory Paper

Total: 4 Credits

Workload :-15 hrs /credit

(Total Workload :- 4 credits :	x 15 hrs = 60 hrs in semester)
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Credit	Description	References
Credit 1	1. Genomics	1. Alwi Z. B. (2005) The Use of SNPs in Pharmacogenomics Studies. <i>Malays J Med Sci.</i> 12(2):4-
	A. Gene sequencing, conserved	12.
	genes, finding base sequences	2. Butler J. M. (2012) Single Nucleotide Polymorphisms and Applications In: Advanced Topics in
	which form genes	Forensic DNA Typing: Methodology. Academic Press: United States.347-369
	B. Many proteins from one gene,	3. Isenbarger T.A., Carr C.E., Johnson S.S., et al. (2008) The most conserved genome segments for
	alternative gene expression: DNA	life detection on Earth and other planets. Orig Life Evol Biosph. 38(6):517-533.
	imprinting and Epigenetics.	4. Brown TA. (2002) Genomes. 2nd edition. Oxford: Wiley-Liss; Chapter 7, Understanding a
	C. Genomic variation-SNPs, SNPS	Genome Sequence. Available from: https://www.ncbi.nlm.nih.gov/books/NBK21136/
	and diseases, SNPS detection and	5. Stojanovic N., Florea L., Riemer C., Gumucio D., Slightom J., Goodman M., Miller W., and
	medical therapies. Eukaryotic and	Hardison R. (1999) Comparison of five methods for finding conserved sequences in multiple
	prokaryotic SNPs	alignments of gene regulatory regions, Nucleic Acids Research, 27 (19) 1,3899-3910.
	D. Role of genomic variation in	6. Lemaître J. F., Berger V., Bonenfant C., Douhard M., Gamelon M., Plard F. and Gaillard J.M.
	aging, Recognition of trades offs	(2015) Early-late life trade-offs and the evolution of ageing in the wild. Proc Biol Sci. 7;
	associated with genomic variation.	282(1806): 20150209.
		7. Morris B. J., Willcox B. J and Donlon T.A. (2019) Genetic and epigenetic regulation of human
		aging and longevity. Biochim Biophys Acta Mol Basis Dis. 1; 1865(7):1718-1744.
		8. Primrose S. B. and Twyman R. M. (2006) Principles of Gene Manipulation and Genomics, 7th

- Edition. S. B. Primrose & R. M. Twyman. Blackwell Publishing: U.S. 626 pp.
- 9. Ramírez-Bello J. and Jiménez-Morales M. (2017) Functional implications of single nucleotide polymorphisms (SNPs) in protein-coding and non-coding RNA genes in multifactorial diseases. Gac Med Mex. 153(2):238-250.
- 10. Shaw V., Bullock K. And Greenhalf W. (2016) Single-Nucleotide Polymorphism to Associate Cancer Risk. Methods Mol Biol. 1381:93-110.
- 11. Watson J. D., Baker T. A., Gann A., Bell S. P., Levine M. and Losick R. 7th Edition. Molecular Biology of the Gene. Pearson-USA
- 12. Yashin A. I., Ukraintseva S. V., Akushevich I. V., Arbeev K. G., Kulminski A. and Akushevich L. (2009) Trade-off between cancer and aging: what role do other diseases play? Evidence from experimental and human population studies. Mech Ageing Dev. 130(1-2):98-104.
- 13. Kaeberlein M. (2013) Longevity and aging. F1000Prime Rep. 5:5.

Credit 2	2. Genetically modified plants and	1. Cotrim A.P. and Baum B. J. (2008) Gene therapy: some history, applications, problems, and
	animals	prospects. Toxicol Pathol. 36(1): 97-103.
	a) Genetically modified organisms-	2. Gene Therapy Tools and Potential Applications- Francisco Martin Molina (2013) Janeza Trdine
	social and ethical issues	9, 51000 Rijeka, Croatia (online book)
	b) Gene augmentation and gene	3. Glick B. R. and Pasternak J. J. (1998) Molecular Biotechnology: Principles and Applications of
	therapy	Recombinant DNA. Washington D C, ASM Press.
	c) Applications in medicine –	http://library.um.edu.mo/ebooks/b28045804.pdf
	prevention, early detection and	4. Weaver R. (2007) Molecular Biology. 4 th Edition. Mc-Grew Hill Publication
	cure of diseases	5. Worgall S. and R. G. (2014) Gene Therapy In: Principles of Tissue Engineering (Fourth Edition).
	d) Applications of transgenic plants	Academic Press: United States. Chapter 34. 657-686.
	and animals - advantages and	6. Maghari B. M. and Ardekani A.M. (2011) Genetically modified foods and social concerns.
	disadvantages	Avicenna J Med Biotechnol. 3(3):109-17.
		7. Agnès E. Ricroch, Michèle Guillaume-Hofnung and Marcel Kuntz (2018) The ethical concerns
		about transgenic crops. Biochem J 475 (4): 803–811.
		8. Ormandy E.H., Dale J. and Griffin G. (2011) Genetic engineering of animals: ethical issues,
		including welfare concerns. Can Vet J. 52(5):544-50.
Credit 3	3. Mobile DNA elements	1. Lewin B. (2011) Genes X. Jones and Bartlett Publication.
	a) Transposable elements in bacteria,	2. Watson J. D., Baker T. A., Gann A., Bell S. P., Levine M. and Losick R. 7 th Edition.
	IS elements, composite transposons,	Molecular Biology of the Gene. Pearson-USA
	Integrons.	3. Lodish H. F. (2003) Molecular Cell Biology 5 Th Edition. New York: W H and Freeman

	b) Replicative, nonreplicative	Company.
	transposons, and Mu transposition	4. Reddy, A.R., Peterson, P.A. Transposable elements of maize. <i>Molec Gen Genet</i> 192, 21–31
	c) Controlling elements in Tn A, Tn 5	5. Kaminker, J.S., Bergman, C.M., Kronmiller, B. et al. (2002) The transposable elements of
	and Tn 10 transposition	the Drosophila melanogaster euchromatin: a genomics perspective. Genome
	d) Transposons in maize and	Biol 3, research0084.1 (2002).
	Drosophila	6. Krastanova, O, M. Hadzhitodorov & M. Pesheva (2005) Ty Elements of the
	e) Retroviruses and retrotransposon,	Yeast Saccharomyces Cerevisiae, Biotechnology & Biotechnological Equipment, 19:
	Ty elements in yeasts	sup2, 19-26,
	f) SINES, LINES and Alu elements.	7. Griffiths AJF, Gelbart WM, Miller JH, et al. (1999) Modern Genetic Analysis. New York: W.
		H. Freeman; Ty Elements in Yeast. Available from:
		https://www.ncbi.nlm.nih.gov/books/NBK21285/
		8. Carnell A. M. and Goodman J.I. (2003) The Long (LINEs) and the Short (SINEs) of It:
		Altered Methylation as a Precursor to Toxicity. Toxicological Sciences. 75(2):229–235
		9. Konkel M. K., Walker J. A. and Batzer M. A. (2010) LINEs and SINEs of primate evolution.
		Evol Anthropol. 1; 19(6):236-249.
		10. Kramerov D. A. and Vassetzky N. S. (2011) Origin and evolution of SINEs in eukaryotic
		genomes. Heredity (Edinb). 107(6):487-95.
		11. Weiner A. M. (2002) SINEs and LINEs: The art of biting the hand that feeds you. Current
		Opinion in Cell Biology. 14(3): 343-350
Credit 4	4. Proteomics	1. Kellner R. (2000) Proteomics: Concepts and perspectives. Fresenius J Anal Chem. 366(6-
	a) Basic concept of proteomics	7):517-524.

- b) Expression, Analysis and Characterization of Protein.
- c) Analysis of protein structure
- d) Protein interaction.
- e) Basic concept of Metabolomics with examples and global biochemical networks

- 2. Graves P.R. and Haystead T. A. (2002) Molecular biologist's guide to proteomics. Microbiol Mol Biol Rev. 2002 Mar;66(1):39-63.
- 3. Bhushan Patwaradhan and Rathnam Chaguture (2005) An overview of the basics of proteomics. In: Innovative approaches in drug discovery, Academic Press: United States. Link to the pdf: https://analyticalscience.wiley.com/do/10.1002/sepspec.10201education/full/
- 4. Baidoo E. E. K. (2019) Microbial Metabolomics: A General Overview. Methods Mol Biol. 1859:1-8.
- 5. Ekman R., Silberring J., Brinkmalm A. W. and Kraj A. (2009) Mass Spectrometry: Instrumentation, interpretation and applications, John Wiley and Sons. Inc., Canada.
- 6. Nölting B. (2006) Methods in Modern Biophysics. Second Edition, Springer: Germany.
- 7. Tang J. (2011) Microbial metabolomics. Curr Genomics. 12(6):391-403.
- 8. Villas-Bôas S. (2012) Katya Ruggiero Microbial Metabolomics CABI.
- 9. Webster D. (2000) Protein Structure, Prediction methods and Protocols. Methods in Molecular Biology Vol 143 Humana Press.
- 10. Wilson K. And Walker J. (2005) Principles and Techniques of Biochemistry and Molecular Biology, 6th Edn., Cambridge University Press, New York.
- 11. Banaei-Esfahani A, Nicod C, Aebersold R, Collins BC. (2017) Systems proteomics approaches to study bacterial pathogens: application to Mycobacterium tuberculosis. Curr Opin Microbiol. 39:64-72.
- 12. Chen B, Zhang D, Wang X, Ma W, Deng S, Zhang P, Zhu H, Xu N, Liang S. (2017) Proteomics progresses in microbial physiology and clinical antimicrobial therapy. Eur J Clin

	Microbiol Infect Dis.36(3):403-413.
	13. Chen F, Ma R, Chen XL. (2019) Advances of Metabolomics in Fungal Pathogen-Plant
	Interactions. Metabolites. 15;9(8):169.
	14. Zhao J., Wang G., Chu J. and Zhuang Y. (2019) Harnessing microbial metabolomics for
	industrial applications. World J Microbiol Biotechnol. 36(1):1-8.
	15. Ramanathan M., Porter D.F. and Khavari P.A. (2019) Methods to study RNA-protein
	interactions. Nat Methods. 16(3):225-234.
	16. Luger K. and Phillips S.E. (2010) Protein-Nucleic acid interactions. Curr Opin Struct Biol.
	20(1):70-2.

MB CCTP- 9 Clinical Microbiology Core Compulsory Theory Paper

Total: 4 Credits Workload :-15 hrs /credit

(Total Workload: -4 credits x 15 hrs = 60 hrs in semester)

Credit	Credit Title and Content	References
		References
Credit	A Determinants of Microbial Pathogenicity	A Determinants of Microbial Pathogenicity
1	i. Adhesion	1. Gal-Mor B. and Finlay B. B. (2006) Pathogenicity islands: a molecular toolbox for
	ii. Invasion	bacterial virulence. Cellular Microbiology. 8 (11): 1707-1719.
	iii. Evasion	2. Iglewski B. H. (1990) Molecular Basis of Bacterial Pathogenesis, first edition, Academic
	iv. Toxigenesis (mode of action –In vivo and In	Press: United States.
	vitro assay systems for diphtheria, cholera,	3. Kudva I. T., Cornick N. A., Plummer P. J., Zhang Q., T. L., Bannantine J.P. and
	tetanus toxoid and endotoxins of Gram	Bellaire B. H. (2016) Virulence Mechanisms of Bacterial Pathogens. Fifth Edition, ASM:
	negative bacteria)	Washington.
	v. Bacterial resistance to host defenses-	4. Peterson J. W. (1996) Bacterial Pathogenesis In: Medical Microbiology, 4 th Edition.
	Phagocytosis, specific and nonspecific	Editor by Samuel Baron, Galveston, Texas, Link to the book:
	humoral factors)	https://www.ncbi.nlm.nih.gov/books/NBK8526/
	vi. Molecular basis of bacterial pathogenicity –	5. Rosenberg E. (2005) The diversity of bacterial pathogenicity mechanisms. Genome
	Cytoskeletal modulation of host cell.	Biol.6(5):320
	Virulence genes and pathogenicity islands.	6. Schmidt H. and Hensel M. (2004) Pathogenicity islands in bacterial pathogenesis. Clin
	B. Disease Prediction Epidemiological Models:	Microbiol Rev. 17(1):14-56.
	i. Introduction to epidemiological modeling for	B. Disease Prediction Epidemiological Models:
	infectious disease dynamics	1. Hethcote H. W. (1989) The basic epidemiology models: models, expressions for r0,
	ii. Types of Models:	parameter estimation, and applications mathematical understanding of infectious disease
	a. Susceptible infectious recovered (SIR)	dynamics. © World Scientific Publishing Co. Pte. Ltd. 1-61
	b. Susceptible exposed infectious recovered	2. Li L., Yang Z., Dang Z., et al. (2020), Propagation analysis and prediction of the COVID-
	(SEIR)	19. Infect Dis Model, 5: 282-292
	iii.A case study:Disease Prediction	3. Siettos C.I. and Russo L. (2013) Mathematical modeling of infectious disease dynamics.
	Epidemiological Models COVID 19	Virulence. 4(4):295-306.
		4. Wearing H.J., Rohani P.and Keeling M.J. (2005) Appropriate models for the management

		of infectious diseases. PLoS Med 2(7): e174 5. Yang Z., Zeng Z., Wang K., Wong S., <i>et al.</i> , (2020) Modified SEIR and AI prediction of the epidemics trend of COVID-19 in China under public health interventions. Journal of Thoracic Disease. 12(3): 165-174
TC 2	Bacterial diseases with respect to causative agents, general characters, detection methods, therapeutic agents and prophylaxis. Handling and disposing of infectious material a. Helicobacter pylori b. Campylobacter jejuni c. Mycobactertium tuberculosis d. Acinetobacter boumanii e. Actinomycetes bovis/israelli	 Asif M., Alvi I.A. and Rehman S.U. (2018) Insight into <i>Acinetobacter baumannii</i>: pathogenesis, global resistance, mechanisms of resistance, treatment options, and alternative modalities. Infect Drug Resist. 11:1249-1260. Available from: https://www.intechopen.com/books/mycobacterium-research-and-development/virulence-factors-and-pathogenicity-of-mycobacterium tuberculosis infection. Mediterr J Hematol Infect Dis. 16; 5(1):e2013070. Echeverria-Valencia G., Flores-Villalva S.and Espitia C.I. (2017). Virulence Factors and Pathogenicity of <i>Mycobacterium</i>. Chapter 12. Mycobacterium - Research and Development. Editor-Wellman Ribón, IntechOpen. Idowu A., Mzukwa, A., Harrison, U., Palamides P., Haas R., Mbao M., Mamdoo R., Bolon J., Jolaiya T., Smith S., Ally R., Clarke A. and Njom H. (2019) Detection of <i>Helicobacter pylori</i> and its virulence genes (<i>cagA</i>, <i>dupA</i> and <i>vacA</i>) among patients with gastroduodenal diseases in Chris Hani Baragwanath Academic Hospital, South Africa. <i>BMC Gastroenterol</i>. 19:73. Jianjun S., Champion P. A. and Bigi F. (2019) Cellular and Molecular Mechanisms of <i>Mycobacterium tuberculosis</i> Virulence. Frontiers in Cellular and Infection Microbiology. 9:331. Joly-Guillou ML. (2005) Clinical impact and pathogenicity of Acinetobacter. Clin Microbiol Infect. 11(11):868-873. Kao C. Y., Sheu B. S. and Wu J. J. (2006) Helicobacter pylori infection: An overview of bacterial virulence factors and pathogenesis. Biomedical Journal 39, 1, 14-23. Kusters J.G., van Vliet A.H.and Kuipers E. J. (2006) Pathogenesis of Helicobacter pylori infection. Clin Microbiol Rev. 19(3):449-490. Lee C.R., Lee J.H., Park M., Park K.S., Bae I.K., Kim Y.B., Cha C.J., Jeong B.C.and Lee S.H. (2017) Biology of <i>Acinetobacter baumannii</i>: Pathogenesis, Antibiotic Resistance Mechani

TC 3	Viral diseases with respect to causative agents, general characters, detection method, therapeutic	 Levin R. E. (2007) Campylobacter jejuni: A Review of its Characteristics, Pathogenicity, Ecology, Distribution, Subspecies Characterization and Molecular Methods of Detection, Food Biotechnology, 21(4): 271-347. Morris F.C., Dexter C., Kostoulias X., Uddin M.I. and Peleg A.Y. (2019) The Mechanisms of Disease Caused by Acinetobacter baumannii. Front. Microbiol. 10:1601. Nyati K. K. (2013) Role of Campylobacter jejuni Infection in the Pathogenesis of Guillain-Barré Syndrome: An Update. Biomedical Research Journal.1-13. Pine L., Howell A. Jr and Watson S.J. (1960) Studies of the morphological, physiological, and biochemical characters of Actinomyces bovis. J Gen Microbiol. 23:403-424. Ricke S.C., Feye K.M., Chaney W.E., Shi Z., Pavlidis H. and Yang Y. Developments in Rapid Detection Methods for the Detection of Foodborne Campylobacter in the United States. Front Microbiol. 9:3280. Misawa N. and Blaser M.J. (2000) Detection and Characterization of Autoagglutination Activity by Campylobacter jejuni Infection and Immunity. 68(11): 6168-6175. Sharma S., Hashmi M.F. and Valentino III DJ. (2020) Actinomycosis. In: StatPearls [Internet]. Treasure Island (FL): StatPearls. Available from: https://www.ncbi.nlm.nih.gov/books/NBK482151/ Testerman T.L. and Morris J. (2014) Beyond the stomach: an updated view of Helicobacter pylori pathogenesis, diagnosis, and treatment. World J Gastroenterol. 20(36):12781-12808. Wong D., Nielsen T. B., Bonomo R. A., Pantapalangkoor P., Luna B., Spellberg B. (2016) Clinical and Pathophysiological Overview of Acinetobacter Infections: a Century of Challenges Clinical Microbiology Reviews 30(1): 409-447. Chauhan N., Narang J., Pundir S., Singh S. and Pundir C. S. (2012). Laboratory diagnosis of swine flu: A review. Artificial cells, blood substitutes, and immobilization
		Clinical and Pathophysiological Overview of <i>Acinetobacter</i> Infections: a Century of
TC 3	Viral diseases with respect to causative agents.	
	agents and prophylaxis. Handling and disposing of	biotechnology. 41(3): 189-195
	infectious material.	2. Chisari F.V., Isogawa M. and Wieland S.F. (2010) Pathogenesis of Hepatitis B virus
	a. Hepatitis B	infection. Pathol Biol (Paris). 58(4):258-66.
	b. H1N1	3. Falasca L., Agrati C., Petrosillo N., Di Caro A., Capobianchi M.R., Ippolito G. and
	c. HIV	Piacentini M. (2015) Molecular mechanisms of Ebola virus pathogenesis: focus on cell
	d. Oncoviruses	death. Cell Death Differ. 22(8):1250-1259.
	e. Ebola Virus	4. Jilani T.N., Jamil R.T., Siddiqui AH. (2020) H1N1 Influenza (Swine Flu) In: StatPearls
		[Internet]. Treasure Island (FL): StatPearls. Available from:

		https://www.ncbi.nlm.nih.gov/books/NBK513241/
		5. Kawai Y., Kimura Y., Lezhava A, et al. (2012) One-step detection of the 2009
		pandemic influenza A(H1N1) virus by the RT-SmartAmp assay and its clinical
		validation. PLoS One. 7(1):e30236.
		6. Khalafallah M.T., Aboshady O.A., Moawed S.A. and Ramadan M.S. (2017) Ebola virus
		disease: Essential clinical knowledge. Avicenna J Med. 7(3):96-102.
		7. Krajden M., McNabb G. and Petric M. (2005) The laboratory diagnosis of Hepatitis B
		virus. Can J Infect Dis Med Microbiol.16 (2):65-72.
		8. Ravina R., Dalal A, Mohan H., Prasad M. and Pundir C.S. (2020) Detection methods for
		influenza A H1N1 virus with special reference to biosensors: a review. Biosci Rep.
		40(2): BSR20193852
		9. Rewar S., Mirdha D. and Rewar P. (2015) Treatment and Prevention of Pandemic
		H1N1 Influenza. Ann Glob Health. 81(5):645-653. doi:10.1016/j.aogh.2015.08.014.
		10. Simon V., Ho D.D. and Abdool Karim Q. (2006) HIV/AIDS epidemiology,
		pathogenesis, prevention, and treatment. Lancet. 5; 368(9534):489-504.
		11. Sullivan N., Yang Z.Y. and Nabel G. J. (2003) Ebola virus pathogenesis: implications
		for vaccines and therapies. J Virol. 77(18):9733-9737.
		12. Wilkins T., Sams R. and Carpenter M. (2019) Hepatitis B: Screening, Prevention,
		Diagnosis, and Treatment. Am Fam Physician. 99(5):314-323.
		13. Wu C.C., Chen Y.S., Cao L., Chen X.W. and Lu M.J. (2018) Hepatitis B virus
		infection: Defective surface antigen expression and pathogenesis. World J
		Gastroenterol. 21; 24(31):3488-3499.
TC 4	Fungal and protozoal diseases with respect to	1. Elewski B.E. (1998) Onychomycosis: pathogenesis, diagnosis, and management. Clin
	causative agents, general characters, detection	Microbiol Rev. 11(3):415-29.
	methods, therapeutic agents and prophylaxis.	2. Hedayati M.T., Pasqualotto A.C., Warn P.A., Bowyer P. and Denning DW. (2007)
	Handling and disposing of infectious material	Aspergillus flavus: human pathogen, allergen and mycotoxin producer. Microbiology.
	a. Candida albicans	153(Pt 6):1677-1692.
	b. Trichophyton metagrophytes	3. Jabra-Rizk M.A., Kong E.F., Tsui C., Nguyen M. H., Clancy C. J., Fidel P. L., Jr. and
	c. Aspergillus flavus	Noverr M. (2016) <i>Candida albicans</i> Pathogenesis: Fitting within the Host-Microbe
	d. Entamoeba histolytica	Damage Response Framework. Infect Immun. 84(10):2724-2739.
	e. Ascaris lumbricoides	4. Kaufman G., Horwitz B. A., Duek L., Ullman Y. and Berdicevsky I. (2007) Infection
	f. Giardia lamblia	stages of the dermatophyte pathogen <i>Trichophyton</i> : microscopic characterization and
	J. State della controlla	2 mg of the defination of the pathogen 1. venophytom interest opic characterization and

- proteolytic enzymes. *Medical Mycology*. 45(2):149–155.
- 5. Martins N., Ferreira I., Barros L., Silva S. and Henriques M. (2014). Candidiasis: Predisposing Factors, Prevention, Diagnosis and Alternative Treatment. Mycopathologia. 177(5-6): 223-240
- 6. Rudramurthy S.M., Paul RA., Chakrabarti A., Mouton J.W. and Meis J.F. (2019) Invasive Aspergillosis by *Aspergillus flavus*: Epidemiology, Diagnosis, Antifungal Resistance, and Management. J Fungi (Basel). 5(3):55
- 7. Petri W. A., Jr. and Singh U.(1999) Diagnosis and Management of Amebiasis. *Clinical Infectious Diseases*. 29(5):1117–1125.
- 8. Kantor M., Abrantes A., Estevez A, Schiller A., Jose Torrent J., Gascon J., Hernandez R. and Ochner C. (2018) *Entamoeba Histolytica*: Updates in Clinical Manifestation, Pathogenesis, and Vaccine Development. *Can J Gastroenterol Hepatol*. 4601420.
- 9. Scott M. (2008). *Ascaris lumbricoides*: a review of its epidemiology and relationship to other infections. Annales Nestlé (English ed.). 66. 7-22.
- 10. de Lima Corvino D.F. and Horrall S. Ascariasis.(2020) In: StatPearls [Internet]. Treasure Island (FL): StatPearls Available from: https://www.ncbi.nlm.nih.gov/books/NBK430796/
- 11. Rumsey P. and Waseem M. *Giardia Lamblia* Enteritis (2020). In: StatPearls [Internet]. Treasure Island (FL): StatPearls Available from: https://www.ncbi.nlm.nih.gov/books/NBK531495/
- 12. Farthing M.J.G. (1993) Pathogenesis of giardiasis. *Transactions of The Royal Society of Tropical Medicine and Hygiene*. 87(3):17–21.
- 13. Hooshyar H., Rostamkhani P., Arbabi M.and Delavari M. (2019) *Giardia lamblia* infection: review of current diagnostic strategies. Gastroenterol Hepatol Bed Bench 12(1):3-12.

MB TE 31 Cell Culture Techniques

Choice based Optional Theory Paper (Elective)

Total: 2 Credits Workload:-15 hrs /credit

(Total Workload :- 2 credits x 15 hrs = 30 hrs in semester

Credit	Credit Title and Contents	References
No.		
1	 Animal Cell Culture Techniques: A. Definition of terms: Primary cell cultures and cell lines, established cell lines, suspension and anchorage dependent cell cultures. B. Transformation of cells in culture, culture media, factors affecting cells in culture. 	 Freshney R. I. (2005) Culture of Animal Cells: A Manual of Basic Technique.5th Ed. John Wiley and Sons, Inc. Masters J. R. W. (2000). Animal Cell Culture – A Practical Approach. 3rd Ed. Oxford University Press. Mather J. P. and Penelope E. R. (1998) Introduction to Cell and Tissue Culture Theory and Technique. Plenum Press, New York
2	Commonly used cell culture systems and cell lines in immunological studies: A. Cell culture systems and their applications: primary lymphoid cell culture, cloned lymphoid cell lines, hybrid lymphoid cell lines. B. Immuno-modulation	 Kindt T. J., Goldsby R. A., Osborne B. A. and Kuby J. (2007) Kuby Immunology. 6th Ed. W. H. Freeman and Co. Patwardhan B., Diwanay S.and Gautam M. (2006) Botanical immunomodulators and chemoprotectants in cancer therapy. In Drug Discovery and Development Volume I: Drug Discovery. Ed. Chorghade Mukund S. Wiley- Interscience, John Wiley and Sons Inc. USA. 405-424.

MB PE 31 Cell Culture Techniques
Choice based Optional Practical Paper (Elective)

Total: 2 Credits Workload:-30 hrs /credit

(Total Workload :- 2 credits x 30 hrs = 60 hrs in semester

Credit No.	Credit Title and Contents	References
1	 A. Density gradient based separation of peripheral lymphocytes (1) B. Preparation of Lymphocyte culture (1) C. Effect of immunomodulators on lymphocyte proliferation (Stimulatory and inhibitory effect) (2) 	 Freshney R. I. (2005) Culture of Animal Cells: A Manual of Basic Technique, 5th Ed., John Wiley and Sons, Inc. Masters J. R. W. (2000) Animal Cell Culture – A Practical Approach. 3rd Ed. Oxford University Press.
2	A. Chick embryo fibroblast cell culture (1)	 Mather J. P. and Penelope E. R. (1998) Introduction to Cell and Tissue Culture Theory and Technique. Plenum Press, New York Hernandez R. and Brown D.T. (2010) Growth and maintenance of chick embryo fibroblasts (CEF). Curr Protoc Microbiol.17:A.4I.1–A.4I.8

MBTE 32 Bioremediation and Biomass Utilization Choice Based Optional Theory Paper (Elective) Exclusively based on Molecular Biology II CCTP8

Total: 2 Credits

Workload :-15 hrs /credit

(Total Workload :- 2 credits x 15 hrs = 30 hrs in semester

Credit	Credit Title and Contents	References
No. TE1	Bioremediation A. Microbial Degradation of xenobiotics, B. Engineered bio- degradative pathways: Camphor, octane, xylene, naphthalene degradation pathway C. Aromatic compound degradation: Manipulation by plasmid transfer Manipulation by gene alteration	 Glick B. R., Pasternak J. J., Cheryl L. and Patten C. L. (1998) Molecular Biotechnology: Principles and Applications of Recombinant DNA. Washington D C, ASM Press Jaiswal S., Singh D. K. and Shukla P. (2019) Gene Editing and Systems Biology Tools for Pesticide Bioremediation: A Review. Front Microbiol. 10:87 Karpouzas D. G. and Singh B. K. (2006) Microbial degradation of organophosphorus xenobiotics: metabolic pathways and molecular basis. Adv Microb Physiol. 51:119-185. Ramos J. L., González-Pérez M. M. and Caballero A., van Dillewijn P. (2015) Bioremediation of polynitrated aromatic compounds: plants and microbes put up a fight. Curr Opin Biotechnol. 16(3): 275-281. Weaver R. (2007) Molecular Biology. 4th Edition. Mc-Grew Hill Publication.
TE2	Biomass utilization A. Utilization of starch and cellulose; B. Isolation of the prokaryotic and eukaryotic cellulase genes, manipulation of the cellulase gene, advantages of using <i>Zymomonas mobilis</i> C. Alcohol, fructose, and silage production; advantages of each D. Improvisation of the processes of alcohol production	 Glick B. R., Pasternak J. J., Cheryl L. and Patten C. L. (1998) Molecular Biotechnology: Principles and Applications of Recombinant DNA. Washington D C, ASM Press Gupta G. V. (2016) New and Future Developments in Microbial Biotechnology and Bioengineering. <i>Aspergillus</i> System Properties and Applications. Elsevier Book Publication. Lal P.B., Wells F.M., Lyu Y., Ghosh I.N., Landick R. and Kiley P.J. (2019) A markerless method for genome engineering in <i>Zymomonas mobilis</i> ZM4. Front.

	E. F.	Improvisation of the processes of fructose production Commercial production processes of alcohol and fructose	4. 5.	Microbiol. 10: 2216 Sarris, D.and Papanikolaou S. Biotechnological production of ethanol: Biochemistry, processes and technologies. Engineering Life Sciences. 16: 307-329 Weaver R. (2007) Molecular Biology. 4 th Edition. Mc-Grew Hill Publication
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MB PE 32 Bioremediation and Biomass Utilization CBOP-3 Choice Based Optional Practical Paper Exclusively based on Molecular Biology II CCTP8

Total: 2 Credits

(Total Workload :- 2 credits x 30 hrs = 60 hrs in semester

Workload:-30 hrs /credit

Credit No.	Credit Title and Contents	References
1	Bioremediation	1. Arora P. K., Srivastava A., and Singh V. P. (2014) Bacterial degradation of nitrophenols
	1. Degradation of para nitrophenol using	and their derivatives.J Hazard Mater. 266:42-59.
	Pseudomonas putida	2. Bánfalvi G and Antoni F. (1990) DNA-based diagnosis. Orv Hetil. 131(18):953-964.
	2. Low density plastic/bioplastic	3. Kulkarni M. and Chaudhari A. (2006) Biodegradation of p-nitrophenol by <i>P. putida</i> .
	degradation using bacterial isolates	Bioresour Technol. 97(8): 982-988.
	3. Demonstration of DNA finger-printing	4. Kumar Khanna V. (2007) Existing and emerging detection technologies for DNA
	technique	(Deoxyribonucleic Acid) finger printing, sequencing, bio- and analytical chips: a
		multidisciplinary development unifying molecular biology, chemical and electronics engineering. Biotechnol Adv. 25(1):85-98.
		5. Li J., Kim H. R., Lee H. M. and Yu H. C., Jeon E., Lee S. and Kim D. (2020) Rapid
		biodegradation of polyphenylene sulfide plastic beads by <i>Pseudomonas</i> sp. Sci Total Environ. 720:137616.
		6. Qiu X., Wu P., Zhang H., Li M. and Yan Z. (2009) Isolation and characterization of
		Arthrobacter sp. HY2 capable of degrading a high concentration of p-
		nitrophenol. Bioresour Technol. 100(21):5243-5248.
		7. Roohi, Bano K., Kuddus M., Zaheer M.R., Zia Q., Khan MF., Ashraf G.M., Gupta A. and
		Aliev G. (2017) Microbial Enzymatic Degradation of Biodegradable Plastics. Curr Pharm
		Biotechnol. 18(5):429-440.
		8. Sangeetha Devi R., Ramya R., Kannan K., Robert Antony A. and Rajesh Kannan V.

	T	T
		 (2019) Investigation of biodegradation potentials of high density polyethylene degrading marine bacteria isolated from the coastal regions of Tamil Nadu, India Mar Pollut Bull. 138:549-560. 9. Wilkes R. A. and Aristilde L. (2017) Degradation and metabolism of synthetic plastics and associated products by <i>Pseudomonas</i> sp.: capabilities and challenges. J Appl Microbiol. 123(3):582-593.
2	Biomass utilization	1. Larkum A. W., Ross I. L., Kruse O. and Hankamer B. (2012) Selection, breeding and
	1. Biodiesel production using micro-algae	engineering of microalgae for bioenergy and biofuel production. Trends Biotechnol.
	2. Isolation of bio-emulsifier producing	30(4):198-205.
	organisms for degradation of aromatic	2. McGinn P. J., Dickinson K. E., Bhatti S., Frigon J. C., Guiot S. R. and O'Leary S. J.
		· · · · · · · · · · · · · · · · · · ·
	compounds	(2011) Integration of microalgae cultivation with industrial waste remediation for
		biofuel and bioenergy production: opportunities and limitations. Photosynth Res. 109(1-3):231-247.
		3. Muhonja C.N., Makonde H., Magoma G. And Imbuga M. (2018) Biodegradability of
		polyethylene by bacteria and fungi from Dandora dumpsite Nairobi-Kenya. PLoS ONE 13(7): e0198446.
		4. Parmar A., Singh N. K., Pandey A., Gnansounou E. and Madamwar D. (2011)
		Cyanobacteria and microalgae: a positive prospect for biofuels. Bioresour Technol. 102(22):10163-10172.
		5. Viramontes-Ramos S., Cristina Portillo-Ruiz M., Ballinas-Casarrubias Mde L, Torres-
		Muñoz J. V., Rivera-Chavira B. E. and Nevárez-Moorillón G. V. (2010) Selection of
		biosurfactan/bioemulsifier-producing bacteria from hydrocarbon-contaminated
		soil. Braz J Microbiol. 41(3):668-675.

MB TE 33 Microbial Virus Technology

Choice based Optional Theory Paper (Elective)

Workload :-15 hrs /credit

Total: 2 Credits

No.	Topic	Reference
edit	A. Isolation and characterization of bacteriophages i. Abundance of bacteriophages in the environment ii. Bacteriophage Lifecycle-Lytic, Lysogeny and chronic cycle. Genetic basis of lytic and lysogeny cycles B. Isolation of bacteriophages from various environmental samples-(Different methods) River, Intestine, Lakes, Tooth plaque, Ponds, High temp.env. Cockroaches, Raw vegetables, Activated sludge, Fecal matter, Sewage, Soil, Flies, Sewage Treatment plant	 Ahiwale Sangeeta (2013) Bacteriophages against enteric bacterial pathogens and their potential for bioremediation of pathogen infested water bodies. PhD thesis, University of Pune, Pune, Maharashtra Forest Rohwer, Merry Youle, Heather Maughan and Nao Hisakawa (2014) Life in Our Phage World. A centennial field guide to the Earth's most diverse inhabitants. Illustrations by Leah L Pantéa and Benjamin Darby (Book) Hobbs Z. and Abedon S. T. (2016) Virology Diversity of phage infection types and associated terminology: the problem with Lytic or lysogenic. Minireview. FEMS Microbiology Letters, 363, , fnw047 doi: 10.1093/femsle/fnw047, 2016 Ahiwale Sangeeta (2013) Bacteriophages against enteric bacterial pathogens and their potential for bioremediation of pathogen infested water bodies. PhD thesis, University of Pune, Pune, Maharashtra Azeredo J. and Sillankorva S. Editors. (2018) Bacteriophage Therapy from Lab to Clinical Practice. In Methods in Molecular Biology. Walker J. M. Series Editor. Humana Press Book. Springer. Clokie M. R. J. and Kropinski A. M. Editors (2009). Bacteriophages: Methods and Protocols. Volume1: Isolation, Characterization and Interactions. Springer Book
	C. Bacteriophage growth kinetics i. Concept and calculations of EoP, MOI ii. Adsorption rate constant iii. One step growth curve-(Latent peroid, Eclipsed period, Rise period, Plateau, burst size D. Phage based bacterial detection: Phage typing	 Clokie M. R. J. and Kropinski A. M. Editors (2009). Bacteriophages: Methods and Protocols. Volume1: Isolation, Characterization and Interactions. Springer Book Effect of bacterial growth rate on bacteriophage population growth rate, Dominik Nabergoj, Petra Modic, Ales Podgornik, Wiley Microbiology open, 2017 Schofield D.A., Sharp N.J. and Westwater C. (2012) Phage-based platforms for the clinical detection of human bacterial pathogens. Bacteriophage. 2(2):105-283

Credit	A. Bacteriophage as biocontrol agent i. Phage based technology for decontamination of water (drinking water, recreational water, medical waste water)	 Ahiwale Sangeeta (2013) Bacteriophages against enteric bacterial pathogens and their potential for bioremediation of pathogen infested water bodies. PhD thesis, University of Pune, Pune, Maharashtra McLaughlin M. R. and Brooks J. P. (2008) EPA worst case water microcosms for testing phage biocontrol of <i>Salmonella</i>. J Environ Qual. 37: 266-271 Sharma S., Soumya Chatterjee S., Datta S., Rishika Prasad R., Dubey D., Prasad R. K. and Vairale M.G. (2017) Bacteriophages and its applications: an overview. Folia Microbiol. 62(1):17-55 Singh M.K., Maurya A. and Kumar S. (2020) Bioaugmentation for the treatment of waterborne pathogen contamination water. Waterborne Pathogens. 189–203.
	ii. Phage based technology for pathogen control in aqua systems	 Culot A., Grosset N. and Gautier M. (2019) Overcoming the challenges of phage therapy for industrial aquaculture: A review. Aquaculture. Elsevier. 513:734423. Kutter E. and Sulakvelidze A. Editors. (2004) Bacteriophages: Biology and Applications. Edition-illustrated. Publisher-CRC Press. Nakai T. and Park S. C. (2002) Bacteriophage therapy of infectious diseases in aquaculture. Mini-review. Research in Microbiology. 153: 13–18 Vinod M. G., Shiva M.M., Umesha K.R., Rajaveera B.C., Krohne G. and Karunasagar J. (2006) Isolation of Vibrio harveyi bacteriophage with potential for biocontrol of luminous vibriosis in hatchery environments. Aquaculture. 55: 117-124
	iii. Bacteriophages for the biocontrol of biofilms on medical devices	 Ahiwale S.S. (2011) <i>In vitro</i> management of hospital <i>Pseudomonas aeruginosa</i> biofilm using indigenous T7-like lytic phage. Curr. Microbiology. 62:335-340 Haradaa L. K., Silvaa E.C., Camposa W. F., Del Fiola F. S., Vilaa M., Dąbrowskab K., Krylovc V. N. and Balcão V. M. (2018) Applications of bacteriophages: State of the art, Review article. Microbiol Res. 212-213:38-58 Lu T. K. and Collins J. J. (2007) Dispersing biofilms with engineered enzymatic bacteriophage. Proceedings of National Academy of Science. 104: 11197-11202
	Bacteriophage based technology for pathogen control in Poultry	 Żbikowska K, Michalczuk M, Dolka B. (2020) The Use of Bacteriophages in the Poultry Industry. Review. Animals (Basel).10(5):872 Gorski A., Miedzybrodzki R. and Borysowski J. (Editors). (2019) Phage Therapy: A Practical Approach. Springer International Publishing
	B. Bacteriophage Therapy i. Use of bacteriophages as therapeutic agent	 Kutter E. and Sulakvelidze A. Editors. (2005) Bacteriophage Therapy in Humans. Chapter 14. Bacteriophages, biology and applications. CRC Press.

ii. Phage lysine therapy and prohylaxis	 Principi N., Silvestri E. and Esposito S. (2019) Advantages and Limitations of Bacteriophages for the Treatment of Bacterial Infections. Front. Pharmacol. 10: 513 Bacteriophages in Health and Disease Hyman P. and Abedon S. T. Editors. (2012) Bacteriophages in Health and Disease. Volume 24 of Advances in molecular and cellular microbiology. Contributor C.A.B. International. Edition- illustrated. Publisher CABI. Vázquez R., García E. and García P. (2018) Phage lysins for fighting bacterial respiratory infections: a new generation of antimicrobials. Mini review article. Front. Immunol. 9: 2252 Eric E. C. and Adhya S. L. (2015). Phage Therapy: Current Research and Applications. Clinical infectious diseases: an official publication of the Infectious Diseases Society of America. 61(1): 141–142 Gorski A., Miedzybrodzki R. and Borysowski J. (Editors). (2019) Phage Therapy: A Practical Approach. Springer International Publishing
C. Mycoviruses: A new dimension in Microbiology i. Occurrence ii. Taxonomy of Mycoviruses iii. Mycovirus-host interaction mechanisms iv. Characterization Technoiques v. Mycoviruses as biocontrol agents against fungal plant pathogens	 Abid, M., Khan, M., Mushtaq, S., Afzaal, S., and Haider, M. (2018). A comprehensive review on mycoviruses as biological control agent. World Journal of Biology and Biotechnology, 3(2), 187-192. Abbas J. (2016) A Review Paper Mycoviruses. Journal of Plant Pathology and Microbiology. 7 (12): 1-4 Zoll J., Verweij P. E. and Melchers W. J. G. (2018) Discovery and characterization of novel <i>Aspergillus fumigatus</i> mycoviruses. PLoS ONE 13(7): e0200511. Niu Y., Yongze Yuan Y., Mao J., Yang Z., Cao Q., Zhang T., Wang S. and Liu D. (2018) Characterization of two novel mycoviruses from <i>Penicillium digitatum</i> and the related fungicide resistance analysis. Scientific Reports. 8:5513 Kondo H., Chiba S., Toyoda K. and Suzuki N. (2013).Evidence for negative-strand RNA virus infection in fungi. Virology, 435: 201–209
D. Introduction of algal viruses	 Coy S. R., Gann E. R., Pound H. L., Short S. M. and Wilhelm S. W. (2018) Viruses of eukaryotic algae: Diversity, Methods for detection and future directions. Viruses. 10 (9): 487

MB PE 33 Practicals based on Clinical Microbiology and Microbial Virus Technology

Choice based Optional Practical Paper (Elective)

Total: 2 Credits Workload:-30 hrs /credit

(Total Workload :- 2 credits x 30 hrs = 60 hrs in semester

		VORKIOAU: - 2 Credits x 30 nrs = 60 nrs in semester
Credit	Description	References
Credit 1.	A. Collection, Handling, transportation of clinical samples. B. Study of drug resistance pattern for clinical isolates: C. Staphyloccus aureus, Pseudomonas aeruginosa, Candida albicans. D. Microbial assay using combination of antibiotics against resistant species of any bacterial isolate. E. Visit to industry	 Biemer J. J. (1973) Antimicrobial susceptibility testing by the Kirby-Bauer disc diffusion method. Ann Clin Lab Sci. 3(2):135-140. Clinical and Laboratory Standards Institute. (2006). Quality control minimal inhibitory concentration (MIC) limits for broth microdilution and MIC interpretive breakpoints. Supplement M27-S2. Clinical and Laboratory Standards Institute, Wayne, PA. National Committee for Clinical Laboratory Standards (2001) Development of in vitro susceptibility testing criteria and quality control parameters. Approved guideline, 2nd ed. NCCLS document M23-A2. National Committee for Clinical Laboratory Standards, Wayne, PA. National Committee for Clinical Laboratory Standards (2002) Reference method for broth dilution antifungal susceptibility testing of yeasts. Approved standard, 2nd ed. NCCLS document M27-A2. National Committee for Clinical Laboratory Standards, Wayne, PA.

MB PE 33 Practicals based on Clinical Microbiology and Microbial Virus Technology (continued)		
References		
man H.W. (2009) Phage classification and characterization. In: Clokie MRJ, Kropinski Eds) Bacteriophages: methods and protocols, Volume: Isolation, characterization and etions, Vol. 501. Humana Press, New York, ale Sangeeta (2013) Bacteriophagesagainst enteric bacterial pathogens and their ial for bioremediation of pathogen infested water bodies PhD thesis, University of Pune, Maharashtra. Ale S.S. (2011) In vitro management of hospital Pseudomonas aeruginosa biofilm using mous T7-like lytic phage. Curr. Microbiology. 62:335-340 A. and Padilla G. (1997) New thermal inducible phages isolated from tropical soils. It is in Journal of Genetics. 20: 4 (2018). Effect of bacterial growth rate on it is in journal of Genetics. 20: 4 (2018). Effect of bacterial growth rate on it is in journal of Genetics. 20: 4 (2018). Effect of bacterial growth rate on it is in journal of growth rate. Microbiology Open, 7, e00558. E.M. and Elbaz R.M. (2013) Isolation and molecular characterization of three in actinophages specific for Streptomyces flavovirens. Journal of Virology Research. 22-17 Jughlin M.R. and Brooks J.P. (2008) EPA worst case water microcosms for testing phage attrol of Salmonella. J Environ Qual. 37: 266-271 M. G., Shiva M. M., Umesha K. R., Rajaveera B. C., Krohne G. and Karunasagar J. Disolation of Vibrio harveyi bacteriophage with potential for biocontrol of luminous in hatchery environments. Aquaculture. 55: 117-124 J. R., Gann E. R., Pound H. L., Short S. M. and Wilhelm S. W. (2018) Viruses of votic algae: Diversity, Methods for detection and future directions. Viruses. 10: 487. In Jung S. and Williams S. T. (1982) Methods for the direct isolation and enumeration of		

MBCP3: Immunology, Molecular Biology and Clinical Microbiology

Core Compulsory Practical Paper

Total: 4 Credits Workload: -30 hrs /credit (Total Workload: -4 credits x 30 hrs = 120 hrs in semester)

Sr. No.	Description	References	
1.	 Practicals based on CCTP 7 Immunology: Precipitation reactions of Antigen - Antibody: Single radial diffusion. Rocket Immuno - electrophoresis Agglutination techniques: Determination of iso-antibodies titre to human blood group antigens. Demonstration of Western Blotting Visit to institute/industry for demonstration of ELISPOT/CFT/FACS/animal inoculation 	 Axelsen N. H., Kroll J. and Weeke B. (1973) A manual of quantitative immunoelectrophoresis: methods and applications. Scand. J. Immunol. 2(Suppl. 1): 37-46 Galvão de França N.D., Cristovão Poli M.C., Almeida Ramos P.G., Rocha Borsoi C.S. and Colella R. (2011) Titers of ABO antibodies in group O blood donors.Rev Bras Hematol Hemoter. 33: 259–262 Kang S.J., Lim Y.A. and Baik S.Y. (2014) Comparison of ABO antibody titers on the basis of the antibody detection method used. Ann Lab Med. 34:300–306. Laurell C. B. (1966) Quantitative estimation of proteins by electrophoresis in agarose gel containing antibodies. Anal. Biochem. 15:45–52 Vaerman J. P. (1981). Single radial immune diffusion, in methods in enzymology: 73 	
2.	 Practicals based on CCTP8 Molecular Biology II Isolation of Plasmid from Bacteria Study of the process of transformation for the strain improvement Blue white screening/bacterium E. coli using a gene for green fluorescent protein Study of the process of bacterial conjugation and transfer of the gene of interest 	 (Langone, J. J. And Van Vunakis, H, Eds.) New York: 291-305. Green M. R. and Sambrook J. (2018) The Hanahan Method for Preparation and Transformation of Competent <i>Escherichia coli</i>: High-Efficiency Transformation. Cold Spring Harb Protoc. (3):10. Griffiths A. J. F., Miller J. H., Suzuki D. T., et al. (2000) An Introduction to Genetic Analysis. 7th edition. New York: W. H. Freeman; Bacterial conjugation. Available from: https://www.ncbi.nlm.nih.gov/books/NBK21942/ Phornphisutthimas S., Thamchaipenet A. and Panijpan B. (2007) Conjugation in <i>Escherichia coli</i>: A laboratory exercise. Biochem Mol Biol Educ. 35(6):440-445. Sambrook J. and Russell D. (2001) Molecular Cloning: A Laboratory Manual, 3rd edn. Cold Spring Harbor, NY: Cold Spring Harbor Laboratory Press. Wilson K. and Walker J. (2005) Principles and Techniques of Biochemistry and Molecular Biology, 6th Edn., Cambridge University Press, New York. 	
	MBCP3: Immunology, Molecular Biology and Clinical Microbiology (continued)		

Sr. No. Description		References
3.	 A. Isolation and identification of (any 2) 1. Helicobacter pylori 2. Campylobacter jejuni 3. Mycobacterium spegmatis 	 Best C. A. and Best T. J. (2009) Mycobacterium smegmatis infection of the hand. Hand (N Y). 4(2): 165–166. Chon JW, Hyeon JY, Yim JH, et al. (2012) Improvement of modified charcoal-cefoperazone-deoxycholate agar by supplementation with a high concentration of polymyxin B for detection of Campylobacter jejuni and C. coli in chicken carcass rinses. Applied and Environmental Microbiology. 78(5):1624-1626. Ferguson DA Jr, Li C, Patel NR, Mayberry WR, Chi DS, Thomas E. (1993) Isolation of Helicobacter pylori from saliva. J Clin Microbiol. 31(10):2802-2804. Gonsalves CC, Borsoi A, Perdoncini G, Rodrigues LB, do Nascimento VP. (2016) Campylobacter in broiler slaughter samples assessed by direct count on mCCDA and Campy-Cefex agar. Braz J Microbiol. 47(3):764-9. Thomas J.E., Gibson G.R., Darboe M.K., Dale A. and Weaver LT. (1992) Isolation of Helicobacter pylori from human faeces. Lancet. 340(8829):1194-1195. Yamada H., Yamaguchi M., Igarashi Y., Chikamatsu K., Aono A., Murase Y., Morishige Y., Takaki A., Chibana H. and Mitarai S. (2018) Mycolicibacterium smegmatis, basonym Mycobacterium smegmatis, expresses morphological phenotypes much more similar to Escherichia coli than Mycobacterium tuberculosis in quantitative structome analysis and cryoTEM examination. Frontiers in Microbiology. 9: Article 1992 Palange P, Narang R, Kandi V. (2016) Evaluation of Culture Media for Isolation of Mycobacterium Species from Human Clinical Specimens. Cureus. 30; 8(8):e757. Zimhony O., Vilcheze C. and Jacobs W.R.J. (2004) Characterization of Mycobacterium smegmatis expressing the Mycobacterium tuberculosis fatty acid synthase I (fas1) gene. J. Bacteriol. 186, 4051–
	MBCP3	4055 : Immunology, Molecular Biology and Clinical Microbiology (continued)
	B. Isolation and identification of (any2) 1 Candida albicans 2 Trichophyton mentagrophytes 3 Aspergillus flavus.	 Joshi KR, Gavin JB. (1974) A simple laboratory method for the rapid identification of Candida albicans. <i>Pathology</i>. 1974; 6(3):231-233. Meinhof W, Laschka P, Scherwitz C. (1975) A synthetic medium for rapid chlamydospore formation in Candida albicans <i>Mykosen</i>. 18(7):291-298. Gunasekaran M, Hughes WF. (1977) A simple medium for isolation and identification of Candida albicans directly from clinical specimens. <i>Mycopathologia</i>. 61(3):151-157. M. Baxter (1966) Isolation of <i>Trichophyton mentagrophytes</i> from British soil, <i>Sabouraudia</i>, 4, 4,1966,

	207–209.
	5. Sinski JT, Kelley LM, Flynt PM, Miegel J. (1977) Dermatophyte isolation media: quantitative appraisal
	using skin scales infected with Trichophyton mentagrophytes and Trichophyton rubrum. J Clin
	Microbiol. 5(1):34-8.
	6. Taber RA, Schroeder HW. (1967) Aflatoxin-producing potential of isolates of the Aspergillus flavus-
	oryzae group from peanuts (Arachis hypogaea). Appl Microbiol. 15(1):140-144.
C. Viral titration by	1. Alexander D.J. and Chettle N.J. (1977) Procedures for the haemagglutination and the haemagglutination
haemagglutination technique	inhibition tests for avian infectious bronchitis virus. Avian Pathology. 6(1):9-17
(Determination of titre)	2. Costabile M. (2010) Determining the Reactivity and Titre of Serum using a haemagglutination Assay J
	Vis Exp. 2010; (35): 1752. Published online
	3. Noah D.L., Hill H., Hines D., White E.L. and Wolff M.C. 2009 Qualification of the hemagglutination
	inhibition assay in support of pandemic influenza vaccine licensure. Clinical and Vaccine Immunology:
	CVI. 16(4):558-566.
	4. World Health Organization. WHO Collaborating Center for Reference and Research on Influenza
	Chinese National Influenza Center National Institute for Viral Disease Control and Prevention, China
	CDC (2013) Laboratory Procedures. (20 December 2013) Serological detection of avian influenza
	A(H7N9) virus infections by modified horse red blood cells haemagglutination-inhibition assay
D. Demonstration of Cultivation	
of viruses by egg inoculation	Visit to institute/industry for demonstration
technique with pock and	visit to institute, industry for demonstration
plaque detection	

Savitribai Phule Pune Universitry **Syllabus restructuring 2020** M.Sc. Microbiology Part II Semester IV

MB CCTP- 10 Pharmaceutical Microbiology **Core Compulsory Theory Paper**

Workload :-15 hrs /credit **Total: 4 Credits**

(Total Workload :- 4 o		cr	edits x 15 hrs = 60 hrs in semester)
Credit			References
TC1	General introduction to medicinal chemistry		
	A. Definition and explanation of terms used in medicinal	1.	Agarwal S. S. and Paridhavi M. (2007) Herbal drug technology. Universities
	chemistry (HITS, Lead compound, Toxicity studies,		Press (India) Pvt. Ltd
	HTS, ADME).Nomenclature of drugs	2.	Altreuter D. and Clark D. S. (1999) Combinatorial Biocatalysis: Taking the
	B. Historical perspectives, significance of medicinal		Lead From Nature. Curr. Opin. Biotechnol. 10: 130-136
	chemistry	3.	Bentley's Textbook of Pharmaceutics, Ed. E. A. Rawlins, 8th Ed. (2002)
	C. Introduction to modern drug discovery, rational drug		Bailliere Tindall, London
	design, molecular modeling, gene and DNA	4.	Burn J. H. (1957) Principles of Therapeutics. Blackwell Scientific Pub. O. Ltd.
	technology in chemotherapy		Oxford.
	D. Classification of drugs based on therapeutic classes,	5.	Chatwal G. P. (2003) Bio-pharmaceutics and Pharmacokinetics. Himalaya
	target, mechanism of action, chemistry, etc.		Publishing House, Mumbai.
		6.	Committee for the Purpose of Control and Supervision on Experiments on
			Animals (CPCSEA). www.cpcsea.com
		7.	Dewick P. M. (2002). Medicinal natural products: A biosynthetic approach,
			2nd Ed., John Wiley and Sons
		8.	Erhardt P. W. (2006) Medicinal Chemistry in the New Millennium: A Glance
			into the Future, Ed. Chorghade M. S. in Drug discovery and Development
			Volume I: Drug Discovery. Wiley-Interscience, John Wiley and Sons Inc.
			USA. 17-102.
		9.	Graly J. O. and Joubert P.H. (1997) Handbook of Phase I /II clinical drug
			trials, CRC Press
		10	. Iyengar M. A. (1993) Pharmacology of Powdered Crude Drugs. Iyengar

		series. Manipal, India 11. Micheles P. S., Khmelnitsley Y. L., Dordick J. S. and Clark D. S., (1998), Combinatorial Biocatalysis, A Natural Approach to Drug Discovery, Trends in Biotechnol. 16(5): 210-215 12. Satoskar R. S. and Bhandarkar S. D. (1991) Pharmacology and Pharmacotherapeutics, 12th Ed., Vol. 1 and 2. Popular Prakashan, Mumbai. 13. Vyas S. P and Dixit V. R. (2002), Pharmaceutical Biotechnology, CBS Publishers and Distributors, New Delhi
TC2	 Drug development A. Lead optimization: lead likeness, drug likeness, determination of biological, biochemical properties of drug, pharmacovigilance. B. Drug designing: Ligand based receptor based drug design. (Protein Crystallography, molecular docking) C. Drug development: Preclinical development. Toxicity testing – acute, sub acute, chronic. D. Clinical development: Clinical trials (aims, objectives and conduct). Clinical trials I, II, III and IV. 	 Franklin T. J. and Snow G. A. (1975) Biochemistry of Antimicrobial Action. Chapman and Hall, London. 1-22 and 160-174 Gale E. F., Cundliffe E., Reynolds P. E., Richmond M. H. and Waring M. J. (1972) The molecular basis of antibiotic action. John Wiley and Sons. London Goldstein A., Aronow L., and Kalman S. M. (1969) Principles of Drug Action. The Basis of Pharmacology. Harper international edition New York. Lorian V. (1986) Antibiotics in laboratory medicine. 2nd Ed. Williams & Wilkins Publication National Committee for Clinical Laboratory Standards (now Clinical and Laboratory Standards Institute, CLSI). NCCLS: 1997. Methods for dilution antimicrobial susceptibility testing for bacteria that grows aerobically. Approved Standards M7-A4. Villanova, PA: National Committee for Clinical Laboratory Standards (now Clinical and Laboratory Standards Institute, CLSI). NCCLS: 2002.Performance standards for antimicrobial susceptibility testing; 12th information supplement (M100-S1). Villanova, PA;
TC3	Biopharmaceuticals: Regulations and sources A. Regulatory authorities and its role: FDA, WHO and CLSI	Blondelle S. E., Perez-Paya E. and Houghten R. A. (1996) Synthetic Combinatorial Libraries: Novel Discovery Strategy for Identification of Antimicrobial Agents. Antimicrobial Agents and Chemotherapy. 1067–1071

2. Holliger M. A. (2008), Introduction to Pharmacology. 3rd Ed. CRC Press. B. Introduction to pharmacopeia: IP, USP, and BP Taylor and Francis. 3. Indian Pharmacopoeia (IP 2018). 8th Edition. Four Volumes with addendum C. Formulation of following pharmaceutical preparation as per IP: 2019. Published by the Indian Pharmacopoeia Commission (IPC) on behalf Antibiotics (with any one example) of the Government of India, Ministry of Health and Family Welfare. Antipyretics (with any one example) ii. 4. Kokate C. K., Purohit A. P., Gokhale A. B. (2000) Pharmacology, 4th Ed., iii. Steroids (with any one example) Nirali Prakashan. iv. Injectables (Distilled water, Saline) 5. Micheles P. S., Khmelnitsley Y. L., Dordick J. S. and Clark D. S., (1998), Vitamins (with any one example). Combinatorial Biocatalysis, A Natural Approach to Drug Discovery, Trends v. in Biotechnol. 16(5): 210-215 6. Osol A. (1980) Remington's Pharmaceutical Sciences, 16th Ed., Easton, Pennsylvania: Mack Publishing Company. 7. Satoskar R. S. and S. D. Bhandarkar (1991) Pharmacology and Pharmacotherapeutics, 12th Edition. Vol. 1 and 2. Popular Prakashan, Mumbai. 8. Vyas S. P. and Dixit V. R. (2002), Pharmaceutical Biotechnology, CBS Publishers and Distributors, New Delhi 9. Walsh G. (2006). Biopharmaceuticals: Biochemistry and Biotechnology. 2nd edition. Wiley (E-Book, 2013). TC4 Physicochemical properties of drug and drug metabolism A. Passage of molecules through biological barriers. 1. Holliger M. A. (2008) Introduction to Pharmacology. 3rd Ed. CRC Press. Membrane transport (paracellular, transcellular). Taylor and Francis. B. Drug absorption: 2. Kokate C. K., Purohit A. P., Gokhale A. B. (2000) Pharmacology. 4th Ed. Drug dosages, from gastric emptying to gastric Nirali Prakashan. permeability to drug, first pass effect, 3. Micheles P. S., Khmelnitsley Y. L., Dordick J. S. and Clark D. S. (1998) bioavailablity. Combinatorial Biocatalysis. A Natural Approach to Drug Discovery. Trends C. Drug distribution: in Biotechnol. 16(5): 210-215 Drug-plasma/ serum binding, blood brain barrier, accumulations in tissues. D. Drug elimination

Drug excretion, Drug biotransformation,	
Biotransformation reactions, Functionalization,	
Conjugation reaction, Reactions leading to toxic	
metabolites	

MB CCTP 11 Microbial Technology Core Compulsory Theory Paper

Total: 4 Credits (Total Workload :- 4 credits x 15 hrs = 60 hrs in semester)

Credit Title and Contents

A. Designing of bioreactors Design aspects CSTRs:

operational aspects such as working volume,

B. The configuration (placement) of impellers in a

C. Immobilized cell reactors and air-lift reactors—

D. Batch, Fed-batch and Continuous operation:

Applications, advantages and limitations of each

The dimensional ratios of the outer shell, and the

turbines and propellers, and their combinations)

Bioreactor design and operation

baffles and impellers.

Design and operation.

type.

Credit

No.

TC 1

References 1. Bioreactor Design and Product Yield (1992), BIOTOL series, Butterworths Heinemann. 2. Doran P. M. (1995) Bioprocess Engineering Principles. Imprint-Academic Press. Copyright-Elsevier. 3. Lydersen B. K., D'Elia N. A. and Nelson K. M. (Eds.) (1993) Bioprocess Engineering: Systems, Equipment and Facilities. JohnWiley and Sons Inc. vessel and the different types of impellers (types of 4. Maiti B. R. (2018) Principles of Bioreactor Design. Publisher: Viva books 5. McDuffie N. G.(1991) Bioreactor Design Fundamentals 1st Edition, Elsevier: eBook ISBN: 9781483221083 6. Ratledge C. and Kristiansen B. eds. (2001) Basic Biotechnology. 2nd Ed. Cambridge Univ. Press. Cambridge 7. Singh L., Mahapatra D. and Yousuf A. (2019). Bioreactors: Sustainable Design and Industrial Applications in mitigation of GHG emissions. Elsevier. ISBN-0128212640, 9780128212646

Workload :-15 hrs /credit

TC 2 Process Variables and Monitoring A. Process Variables:

- i. Aeration Theory of oxygen transfer in bubble aeration, Oxygen transfer kinetics (Oxygen Uptake Rate –OUR; Oxygen Transfer Rate OTR; Ccrit), determination of KLa.
- ii. Agitation Functions of agitation. Flow patterns with different types of impellers.
 - a) Fermentation broth rheology and power requirements for agitation – Concept of Newtonian and non Newtonian fluids,
 - b) Effect of broth rheology on heat, nutrient and oxygen transfer,
 - c) Reynold's number, Power number, Aeration number: working out examples using different software.

B. Monitoring of processvariables:

- i. Use of various types of sensors and biosensors for monitoring environmental parameters (pressure, pH, temperature, DO and DCO2)
- ii. Basic principles of operation, types of biosensors

- 1. Aiba S., Humphrey A. E. and Millis N. F. (1982). Biochemical Engineering. Second Edition. Academic Press.
- 2. Angela Jozala (2017) Fermentation Processes Publisher-BoD. Books on Demand. ISBN-9535129279, E-Book 9789535129271
- 3. Carl-Fredrik Mandenius. (2016) Bioreactors: Design, Operation and Novel Applications. Reprint. Publisher-John Wiley & Sons. ISBN 3527683372 E-Book- 9783527683376
- 4. Chand Subhash (1998): Fermentation Biotechnology: Industrial Perspectives. Industrial Perspectives: Proceedings of the Symposium on Biotech Industry a Challenge for 2005 A.D. -with Special Reference to Fermentations. November 4-6, 1998. Publisher: All India Biotech Association
- 5. Larroche C., Sanroman M., Du G. and Pandey A. (Editors). (2016) Current Developments in Biotechnology and Bioengineering: Bioprocesses, Bioreactors and Controls. Publisher-Elsevier, ISBN 0444636749, E- Book-9780444636744
- 6. Lydersen B. K., D' Elia N. A. and Nelson K. M. (Eds.) (1993) Bioprocess Engineering: Systems, Equipment and Facilities. John Wiley and Sons Inc.
- 7. Operational Modes of Bioreactors (1992) BIOTOL series, Butter worths Heinemann.
- 8. Stanbury P., Whitaker A. and Hall S. (2016) Principles of Fermentation Technology. 3rd Edition Imprint: Butterworth-Heinemann

TC 3.	Microbial Fermentation Processes: i. Upstream, Fermentation and Downstream Processing for the following:	Arora D. K. (2005) Fungal Biotechnology in Agricultural, Food and Environmental Applications (Mycology), Marcel Dekker, Inc. New York. Basel
	a. Antibiotics (Rifamycin)ii. Microbial enzymes (Chitinase).	2. Belter P.A., Cussler E. L. and Hu W.S. (1994) Bioseparations Downstream processing for Biotechnology. John Wiley and Sons. N.Y. ISBN: 978-0-471-12113-8
	iii. Exopolysaccharides (Pullulan)iv. Use of immobilized cells / enzymes for	3. Crueger W. and Crueger A (1990) Biotechnology: A textbook of Industrial Microbiology. 2nd edition. Sinauer associates, Inc
	hioconversion	4. Klegerman M. E. and Groves M. J. (1992) Pharmaceutical Biotechnology: Fundamentals and Essentials. Interpharm Press Ltd. Buffalo Grove, Illinois
	environmental applications	5. Meshram S. U. and Shinde G. B. (2009) Applied Biotechnology. I.K. International Pvt. Ltd.
		6. Mishra C. S. K., Ed., Pascale Champagne Associate editor. (2009) Biotechnology applications .I. K. International Pvt. Ltd.
		7. Peppler H. J. and Perlman D. (1970) Microbial Technology Volume 1 and 2, Academic Press, New York.
		8. Ponkhshe S. (1988) Management of Intellectual Property, Bhate and Ponkhshe Prakasham, Pune
		9. Reed G. Ed. Prescott and Dunn's Industrial Microbiology. 4th Ed., CBS Pub. New Delhi.
		10. Van Damme E. J. (1984) Biotechnology of Industrial Antibiotics. Marcel

		Dekker Inc., New York.
		11. Wiseman A. (1985) Topics in Enzyme and Fermentation Biotechnology. Vol. 1 and 2. John Wiley and Sons, New York
TC 4.	Principle concepts of IPR, ISO and Validation Process: A. Intellectual Property Rights (IPR): i. Basic concepts of IPR ii. Introduction to forms of IPR – Patents and Designs B. The concept of ISO Certification. C. Preparation of SOPs D. Validation protocols for methods in: i. Quality Control ii. Process validation	 Calnan N., Redmond A. and O'Neill S. (2009). The FDA's draft process validation Guidance A perspective from industry. Process Validation Guidance. Pharmaceutical Engineering. GMP Publishing. 7(4): 1-17 Supplementary Training Modules on Good Manufacturing Practice. Validation WHO Technical Report Series, No.937, 2006, Annex 4.
	The above should be discussed within WHO Norms. Exercises on preparation of SOPs, operation and validation for analytical methods	

MBTE 41 Quality Assurance and Validation in Pharmaceutical Industry and Development of Anti-infectives from plants

Choice based Optional Practical Paper (Elective)

Total: 2 Credits

Workload:-15 hrs/credit

(Total Workload :- 2 credits x 15 hrs = 30 hrs in semester

Credit	Description	References
Credit 1.	Quality Assurance and Validation in	1. Kokate C. K., Purohit A. P. and Gokhale A. B. 2000. Pharmacology, 4th Ed. Nirali
	Pharmaceutical Industry	Prakashan.
	A. Good Manufacturing Practices (GMP) and	2. Holliger M. A. 2008. Introduction to Pharmacology, Third Ed., CRC Press. ISBN
	Good Laboratory Practices (GLP) in	9781420047417
	pharmaceutical industry.	3. Maron D. M. and Bruce N. A. 1983. Revised methods for the Salmonella mutagenicity
	B. Quality assurance and quality management	test. Mutation Research. 113:173-215
	in pharmaceuticals ISO, WHO and US	4. Osol A. and Hoover J. E. 1975. Remington's Pharmaceutical Sciences, 15th Ed., Mack
	certification. Safety in microbiology laboratory.	Pub. Co., Pennsylvania.
	C. Safety profile of drugs:	5. Blondelle S. E., Pérez-Payá E. and Houghten R. A. 1996. Synthetic combinatorial
	i. Strerility Testing	libraries: novel discovery strategy for identification of antimicrobial agents.
	ii. Pyrogenicity testing	Antimicrobial Agents and Chemotherapy, 1067–1071
	iii. Mutagenicity and Carcinogenicity testing	6. Vyas S. P and Dixit V. R. 2002. Pharmaceutical Biotechnology, CBS Publishers and
	iv. Teratogenicity testing	Distributors, New Delhi
Credit 2.	Development of Anti-infectives:	1. Franklin T. J. and Snow G. A., (1975), Biochemistry of Antimicrobial Action,
	Therapeutic ratio, MIC and MBC	Chapman and Hall, London, 1-22 and 160- 174 2.
	Susceptibility Testing:	2. Gale E. F., Cundliffe E., Reynolds P. E., Richmond M. H. and Waring M. J., (1972),
	A. Use of liquid and solid media	The molecular basis of antibiotic action, John Wiley and Sons, London
	B. Factors affecting susceptibility testing,	3. Goldstein A., Aronow L., and Kalman S. M. (1969) Principles of Drug Action, The
	CLSI guidelines	Basis of Pharmacology, Harper international edition New York.
	C. Diffusion methods – agar dilution	4. Lorian V., (1986), Antibiotics in laboratory medicine, 2nd Ed, Williams & Wilkins
	technique, gradient plate techniques,	Publication
	E-test, Kirby Bauer, Stokes method	5. National Committee for Clinical Laboratory Standards (now Clinical and Laboratory
	D. Susceptibility testing for:	Standards Institute, CLSI). NCCLS: 1997. Methods for dilution antimicrobial
	i. Anti-mycobacterial agents	susceptibility testing for bacteria that grows aerobically. Approved Standards M7-A4.
	ii. Anti-fungal agents	Villanova, PA.
	iii. Anti-protozoan agents	6. National Committee for Clinical Laboratory Standards (now Clinical and Laboratory
	iv. Anti-viral agents	Standards Institute, CLSI). NCCLS: 2002. Performance standards for antimicrobial susceptibility testing; 12th information supplement (M100-S1). Villanova, PA

MBPE 41 Practicals based on Pharmaceutical Microbiology

Choice based Optional Practical Paper(Elective)

Total: 2 Credits	Choice based	Workload: -30 hrs /credit
	(Total Workload :-	-2 credits x 30 hrs = 60 hrs in semester
Credit	Description	References
Credit 1.	Sterility testing of following pharmaceutical preparations as per IP: i. Oral preparations preparation: Antipyretic or antibiotic tablets ii.Liquid preparation: water soluble vitamin or cough syrup or ophthalmic drops iii. Bulk preparation: (any two) Surgical Cotton rolls/ gauze/ surgical sutures/ disposable syringes.	 Holliger M. A. (2008) Introduction to pharmacology. 3rd EdCRC Press 38 Indian Pharmacopoeia. (2007) Government of India, Ministry of Health and Family Welfare. The Indian Pharmacopoeia commission. Ghaziabad. 1:53 Knudsen L. F. (1949) Sample size of parenteral solutions for sterility testing. J Amer Pharm Assoc. 38: 332–337. McGuire J. and T.C. Kupiec (2007) Quality-control analytical methods: the quality of sterility testing. Intl J Pharm Compounding 11(1): 52–55. Madsen R. E. (1994) US vs. Barr Laboratories: a technical perspective. PDA J Pharm Sci Tech. 48(4): 176–179. Moldenhauer J. and Sutton S.V.W. (2004). Towards an improved sterility test. PDA J Pharm Sci Tech. 58 (6): 284–286. Moldenhauer J. (2006). Viability-based rapid microbiological methods for sterility testing and the need for identification of contamination. PDA J Pharm Sci Tech. 60(2): 81–88. Schroeder H.G. (2005). Sterility failure analysis. PDA J Pharm Sci Tech. 59(2): 89–95. Sykes G. (1956) The technique of sterility testing. J Pharm Pharmacol.8: 573.
Credit 2.	Detection and isolation of anti- infectives from plant i. Extraction of bioactive principles from plant and activity fractionation ii. Estimation of its antimicrobial activity using standard guidelines (CLSI)	 Lorian V. (1986) Antibiotics in laboratory medicine. 2nd Ed. Williams and Wilkins Publication National Committee for Clinical Laboratory Standards (now Clinical and Laboratory Standards Institute, CLSI). NCCLS: 1997. Methods for dilution antimicrobial susceptibility testing for bacteria that grows aerobically. Approved Standards M7-A4. Villanova, PA. National Committee for Clinical Laboratory Standards (now Clinical and Laboratory Standards Institute, CLSI). NCCLS: 2002. Performance standards for antimicrobial susceptibility testing; 12th information supplement (M100-S1). Villanova, PA.

MBTE 42 - Advances in Microbial Technology

Choice based Optional Theory Paper (Elective)

Total: 2 Credits Workload:-15hrs/credit

(Total Workload :- 2 credits x 15hrs = 30 hrs in semester

	· · · · · · · · · · · · · · · · · · ·	
Credit	Credit Title and Contents	References
No.		
Credit 1	 Microbial Growth characteristics and product formation i. Concept of primary (growth associated) and secondary (growth on associated) metabolites and their control, ii. Kinetics of growth and product formation (growth rate, yield coefficient, efficiency etc.) iii. Effect of type of growth on fermentation: The type of growth (mycelia pellet form, mycelia filamentous form, free cell, cells producing exopolysaccharides) affects mass transfer of nutrients, oxygen and heat; as also cell proliferation can be affected by shearing of cells. At least one example of each type may be explained to show these effects in any suitable fermentation. 	 Dubasi Govardhana Rao, (2010)Introduction to Biochemical Engineering. Tata Mcgraw Hill Education Stanbury P.F. (2009) Principles of Fermentation Technology. 2 Ed, Elsevier (A Division of Reed Elsevier India Pvt. Limited). Vijai Kumar Gupta, Monika Schmoll, Minna Maki, Maria Tuohy, Marcio Antonio Mazutt editors. (2013) Applications of Microbial Engineering. CRC Press
Credit 2	Animal cell culture technology to produce: i. Recombinant forms of natural proteins (insulin, erythropoietin), ii. Recombinant vaccines (protein: HIV, hepatitis B and	 Moo Young M. ed. (1985) Comprehensive Biotechnology Vol: III and IV, Pergamon Press. N. Y Ratledge Cand Kristiansen Beds. (2001) Basic Biotechnology 2nd Ed. Cambridge Univ. Press. Cambridge

DNA: HIV, malaria), Recombinant enzymes(lipase, restriction endonuclease),	3. Satyanarayana U. (2005) Biotechnology. Books and Allied (p) limited.
iii. Monoclonal antibodies	
iv. Nucleic acid based products (introduction to gene therapy)	

MBPE 42 - Advances in Microbial Technology

Choice based Optional Practical Paper(Elective)

Total: 2 Credits Workload: -30 hrs /credit

Total: 2 Credits		Workload: -30 hrs /credit			
	(Total Workload :- 2 credits x 30 hrs = 60 hrs in semester				
Cre	Credit Title and Contents	References			
dit					
1	A. Bioconversion	A. Bioconversion:			
	Bioconversions using	1. Arana-Peña S., Rios N. S., Carballares D., Mendez-Sanchez C., Lokha Y., Gonçalves L. and Fernandez-			
	immobilized systems	Lafuente R. (2020). Effects of enzyme loading and immobilization conditions on the catalytic features of			
	(cells / enzyme) lipase from <i>Pseudomonas fluorescens</i> immobilized on octyl-agarose beads. Frontiers in bioengineering				
	Parameter testing:	and biotechnology. 8:36.			
	a. Effect of gel concentrationb. Effect of cell / enzyme	2.Brena B, González-Pombo P and Batista-Viera F. (2013) Immobilization of enzymes: a literature survey. Methods Mol Biol. 1051:15-31.			
	concentration	3.Gedam P.S., Raut A.N. and Dhamole P.B. (2019). Effect of operating conditions and immobilization on butanol enhancement in an extractive fermentation using non-ionic surfactant. Appl Biochem Biotechnol 187:1424–1436			
	4.Mahajan R., Gupta V.K. and Sharma J. (2010) Comparison and suitability of gel matrix for entrapped and suitability of gel matrix for entrapped and suitability of gel matrix.				
	higher content of enzymes for commercial applications. Indian J Pharm Sci. 72(2):223-228.				
		B. Laboratory scale production			
	B. Laboratory scale	1. Biswas J. and PaulA. K. (2017). Optimization of factors influencing exopolysaccharide production by			
	production	Halomonas xianhensis SUR308 under batch culture. AIMS Microbiology, 3(3), 564–579.			
	Laboratory scale production and	2. Hereher F., El-fallal A. and Abou-Dobara M. (2018). Cultural optimization of a new exopolysaccharide			

media optimization for:	producer "Micrococcus roseus". Beni-Suef University Journal of Basic and Applied Sciences. 7(4): 632-
exopolysaccharide / bioemulsifier	639
production	 Maia P., Santos V., Ferreira A., Luna M., Silva T., Andrade R. and Campos T. G. (2018). An efficient bioemulsifier-producing <i>Bacillus subtilis</i> UCP 0146 isolated from mangrove sediments. Colloids and Interfaces. 2. 58. 10.3390/colloids2040058 Rosero Neira-Gladys; Pimienta Astrid-Lorely.; Dugarte F. and Carvajal Fredy-Gonzalo. (2003) Parameters examination of a biosurfactant production at laboratory scale. C.T.F Cienc. Tecnol. Futuro [online]. 2(4): 35-42
2 Animal Cell Culture	Pandey S. (2010) Hybridoma technology for production of monoclonal antibodies. Pharmaceutical
Technology	Sciences Review and Research. 1(2): Article 017. 88-94
 A. Preparation of Hybridoma from tumour cell lines. B. B. Production of monoclonal antibodies from hybridoma of tumour cell lines 	Carvalho L. S., da Silva O. B., de Almeida G. C., de Oliveira J.D., Parachin N. S. and Carmo T. S. (2017). Production Processes for Monoclonal Antibodies. Fermentation Processes, Angela Faustino Jozala. IntechOpen. Chapter 10.181-198 Kavyasudha C., Joel J. P. and Devi A. (2018) Differential expression of nucleostemin in the cytoplasm and nuclei of normal and cancerous cell lines. Turk J Biol. 42: 250-258 Greenfield E. A. (2014) Generating Monoclonal Antibodies. Chapter 7. Antibodies: A laboratory Manual. 2 nd edition. Cold Spring Harbour Laboratory Press. New York.629-644

MBTE 43: Industrial waste water treatment and Industrial production of vaccines Choice based Optional Practical Paper (Elective)

Workload :-15 hrs /credit

Total: 2 Credits (Total Workload :- 2 credits x 15hrs = 30 hrs in semester

Credit	Description	References
Credit	Concept and Introduction to Primary, Secondary and Tertiary	1. Abdallh M. N., Abdelhalim W. S. and Abdelhalim H. S. (2016) Industrial
I	treatment of Wastewater. (1 Lecture)	wastewater treatment of food industry using best techniques. International Journal
	Biological Treatment- Aerobic and Anaerobic, Suspended and	of Engineering Science Invention, 5(8):15-28.
	Attached growth processes.	2. Ali, Z. and Rahman, M. (2008) Physico-chemical characteristics of pulp and paper
	Activated Sludge treatment and analysis (reactions and	mill effluent. Research in Environment and Life Sciences.1 (2):59-60.
	Kinetics, mass balance analysis, Hydraulic characters) Critical	3. Ashtekar S., Bhandari V.M., Shirsath S.R., Sai Chandra P.L.V.N. and Jolhe P.D.
	Operating parameters like DO, Hydraulic retention time, Mean	(2013) Dye wastewater treatment: removal of reactive dyes using inorganic and
	cell retention time, F/M ratio. (4 or 5 Lectures)	organic coagulants. Journal of Industrial Pollution Control, 30(1):33-42
	Current industrial wastewater treatment processes:	4. Bajpai P. and Bajpai P.K. 1994. Mini review: Biological colour removal of pulp
	Composition, physico-chemical properties and various effluents	and paper mill wastewaters. Journal of Biotechnology. 33: 211-220.
	treatment methods with reference to:	
	A. Dairies (2 Lectures) B. Food processing (2 Lectures)	
	C. Dyeing industry / Dye-house effluents (2 Lectures)	Advances in Applied Microbiology.48:79-134.
		6. Catalkaya E.C. and Kargi F. 2006. Color, TOC and AOX removals from pulp mill
	D. Paper and pulp industry (2 Lectures) Effluent Dispessal and Payer (1 Lecture)	effluent by advanced oxidation processes: A Comparative Study. Journal of
	Effluent Disposal and Reuse (1 Lecture)	Hazardous Materials. 139 (2): 244-253
		7. Metcalf and Eddy (Eds.) 3 rd Edition, Tata Mac Graw Hill Publishing Co. Ltd. New
		Delhi.
		8. Patwardhan A. D. 2008. Industrial wastewater treatment. © Prentice – Hall of

		India Pvt. Ltd., New Delhi. ISBN 978-81-203-335
		9. Tchobanoglous G. and Burton F. L. (1991) Wastewater engineering, treatment,
		disposal and reuse. 3 rd Edition, Metcalf and Eddy (Eds.), Tata Mac Graw Hill
		Publishing Co. Ltd. New Delhi.
Credit	Industrial production of vaccines	
2	A. Introduction to vaccines	1. Casida, L. E. (1984) Industrial Microbiology. Wiley Easterbs, New Delhi
	B. Types:	2. Patel A. H. (1985) Industrial Microbiology, Macmillan India Ltd.
	In activated, Attenuated, Toxoid, Subunit, Conjugate, Experime	3. Soma Marla S., Bonthala V. S., München H. Z., Suresh., Gaur V. S. and Gohar Taj G.
	ntal,	(2012) Biotechnology in Medicine and Agriculture Principles and Practices. Publisher: I.K
	Valence, Heterotypic	International Publishing House pvt.ltd, Editors: Anil Kumar, Ashwani Pareek, Sanjay
	C. Production	Mohan Gupta. 739-759
	a. Pilot and Industrial scale production	4. Stanbury P. F. and Whittaker A. (1984) Principles of Fermentation Technology.
	b. Excipients	Pergamon press.
	c. Role of Adjuvants and preservatives	5. https://www.slideshare.net/adammbbs/pathogenesis-3-rd-internal-updated-43458567
	D. Production of viral, bacterial and protozoal vaccines -	6. https://www.bio.fiocruz.br/en/images/stories/pdfs/mpti/2013/selecao/vaccine-process-
	Generations of vaccines:	technology.pdf
	i. First generation vaccines-Live attenuated (BCG, MMR)	7. https://www.dcvmn.org/IMG/pdf/ge_healthcare_dcvmn_introduction_to_pd_for_vaccine_
	and Inactivated (Pertussis, Tetanus toxoids)	production_29256323aa_10mar2017.pdf
	ii. Second generation vaccines(synthetic)	8. https://www.sciencedirect.com/science/article/pii/B9780128021743000059
	protein/peptide/polysaccharide) —	9. https://www.researchgate.net/publication/313470959_Vaccine_Scale-
	a. Subunit vaccines (HepB)	up_and_Manufacturing
	b. Recombinant (Rotavirus),	
	c. Hapten-Conjugate vaccines (diphtheria)	

iii.	Third generation vaccines – DNA/RNA and Idiotype
	vaccines (Malaria)
iv.	Next generation vaccines using OMICs approach:
	SARS.

MBPE 43 Practicals based on Industrial Waste Water Treatment and Industrial Production of Vaccines

Choice based Optional Practical Paper(Elective)

Total: 2 Credits Workload: -30 hrs /credit

(Total Workload :- 2 credits x 30 hrs = 60 hrs in semester References Credit Description 1. Barthwal R. R. (2002) Environmental Impact Assessment, New Delhi (India). New Age i. Estimation of pollution load of Credit 1. International (P) Limited Publishers. **Practicals** a natural sample (e.g. river water / industrial waste water) 2. Eaton A. D. (2005) Standard methods for the examination of water and wastewater. American based on Public Health Association. American Water Works Association. Water Environment Federation. industrial ii. Setting up a laboratory experiment to assess Publisher: Washington, D.C.: APHA-AWWA-WEF. National government publication: English: waste water degradability of synthetic waste 21st edition treatment 3. Glasson J., Therivel R. and Chadwick A. (2012) Rutledge-Taylor and Francis Introduction to water Environmental Impact Assessment. 4th Edition. 416 pages 4. Srivastava A.K. (2003) Environment Impact Assessment, (A.P.H. Publishing. Corporation, Delhi, ISBN-817648-4423, Credit 2. 1. Cruickshank R. (1982) Medical Microbiology, 12th Edition, P.403. i. Checking the potency of a **Practicals** toxoid based vaccine by 2. Felix A. (1942) Brit. Med. J. 11: 597. immune diffusion assay 3. Roitt L. (1994) Essential Immunology. 8th edition. Blackwell Scientific. Oxford, UK.114-115. based on industrial ii. Preparation of Salmonella O 4. Vaerman J.P. (1981) Single radial immune diffusion, in methods in enzymology. 73 (Langone, J. J. and H antigen and estimation And Van Vunakis, H. Eds.) New York. 291-305. production of vaccines with known antibodies

MBTE 44 Bioethics, Biosafety, Quality Control and Quality Assurance Choice based Optional Theory Paper (Elective)

Total: 2 Credits

Workload :-15 hrs /credit

	(Total Workload :- 2 credits x 150 l	ars = 30 hrs in semester
Credit		References
No.		
1	Bioethics and Biosafety A. Bioethics i. Concept of ethics and bioethics with respect to microbiological research ii. Principles of bioethics. iii. Ethical conflicts in microbiological and biotechnological research iv. Biological Diversity Act: conservation of biological diversity, sustainable use of its components and fair and equitable sharing of the benefits arising out of utilization of genetic resources B. Biosafety Regulatory bodies (Role and functions) i. Advisory Committee: Recombinant DNA Advisory Committee (RDAC) ii. Regulatory / Approval Committees: a. Genetic Engineering Appraisal Committee (GEAC) b. Review Committee on Genetic Manipulation (RCGM) c. SIRO (DSIR) d. Institutional Biosafety Committee (IBSC): Importance of Biosafety Institutional Biosafety Committees (IBSCs) Laboratory associated infections and hazards Bio safety regulation: handling of recombinant DNA products and process in industry and in institutions iii. Monitoring Committees: a. State Biotechnology Coordination Committee (SBCC)	 Biotechnology: A comprehensive treatise (Vol. 12). Legal economic and ethical dimensions VCH. (2nded) ISBN- 10 3527304320. 2. Encyclopedia of Bioethics 5 vol set, (2003) ISBN-10: 0028657748. Thomas, J.A., Fuch, R.L. (2002). Biotechnology and safety Assessment (3rd Ed) Academic press. Notification from Department of Biotechnology, Ministry of Science and Technology, India. (2020) Revised simplified procedures/guidelines on Import, Export and Exchange of GE organisms and product thereof for R& D purpose. File no. BT/BS/17/635/2015-PID. dated-17/01/2020 https://ibkp.dbtindia.gov.in/ Ministry of Law And Justice (Legislative Department) New Delhi, the 5th February, 2003/Magha 16, 1924 (Saka) published for general information: The Biological Diversity Act, 2002 No. 18 of 2003 [5th February, 2003]

b	b. District Level Committee (DLC)	
2 Quanta A. B. C. D. i. ii. iiv v. vi	Assessment of suitability of components and products Evaluation of the performance of the manufacturing process Quality Assurance reviewing and approval of procedures, reviewing records and performing audits Good Manufacturing Practices (GMP) and Good Laboratory Practices (GLP) Regulatory bodies (Role and functions): The Central Drugs Standard Control Organization (CDSCO) National Accreditation Board for Testing and Calibration Laboratories (NABL) Food Safety and Standards Authority of India (FSSAI): Food and water Laboratories International Standard ISO/IEC 17025:2017(E). Bureau of Indian Standards -IS 14648 (2011): Methods of Test for Microbiological Examination of Industrial Product (examples Cosmetics And Cosmetic Raw Materials)	 Draft Manual on method of microbiological testing (2016) microbiology of foods. Food safety and Food Standards. Available as https://old.fssai.gov.in/Portals/0/Pdf/Microbiological_Testing_Fo ods_Draft_Manual_06_09_2016.pdf Eleftheriadou M. and Tsimillis K. C. (Eds), Eurachem guide: Accreditation for Microbiological Laboratories, Second edition (2013), ISBN: 978-91-87017-92-6. Available from www.eurachem.org. http://www.electropedia.org/ https://archive.fssai.gov.in/home/food-testing/food-testing-manual.html. https://cdsco.gov.in/opencms/opencms/en/About-us/Functions/ https://cdsco.gov.in/opencms/opencms/en/Home/ https://cpcb.nic.in/functions/ https://www.iso.org/obp International Standard ISO/IEC 17025:2017(E). General requirements for the competence of testing and calibration Laboratories. Third edition. 2017-11 IS 14648 (2011): Methods of Test for Microbiological Examination of Cosmetics And Cosmetic Raw Materials. Available at: https://law.resource.org/pub/in/bis/S11/is.14648.2011.pdf Manual for Good Food Laboratory Practices (GFLPs). 2018. Food Safety and Standards Authority of India (FSSAI), Ministry Of Health and Family Welfare Government Of India, New Delhi Manual of Methods for Analysis of Water 2016. Food Safety and Standards Authority of India (FSSAI), Ministry Of Health and Family Welfare Government of India, New Delhi National Accreditation Board for Testing and Calibration

Laboratories (NABL). (2019)Specific Criteria for Accreditation.
NABL 112. Issue No: 04. Issue Date -11-Feb-2019

	MDDE 44 Dreaticals based on Picothias Picosfety, Quality Control and Quality Assurance				
	MBPE 44 Practicals based on Bioethics, Biosafety, Quality Control and Quality Assurance Choice based Optional Practical Paper (Elective)				
Total	: 2 Credits	Workload :-30 hrs /credit			
	(Total Workload :- 2 cred	lits x $30 \text{ hrs} = 60 \text{ hrs in semester}$			
Sr.	Description	References			
No.					
1.	NABL norms for Calibration of:	National Accreditation Board for Testing and Calibration Laboratories			
	i. Autoclave- Calibration of pressure gauge and temperature by	(NABL). (2019)Specific Criteria for Accreditation. NABL 112. Issue No: 04			
	thermal mapping, sterility testing, SOP preparation.	Issue Date:11-Feb-2019			
	ii. Laminar Air Flow- checking the functioning of UV light by				
	colony count method and sterility checking by blood agar media				
	plate method, SOP preparation.				
2	Food Safety and Standards Authority of India (FSSAI)	Manual of Methods for Analysis of Water 2016. Food Safety and Standards			
	Regulations Test Methods for Drinking Water	Authority of India (FSSAI), Ministry Of Health and Family Welfare			
	i. Detection of sulphite-reducing anaerobes (Clostridia)	Government of India, New Delhi			
	ii. Detection of viruses				
3	Food Safety and Standards Authority of India (FSSAI)	Manual of Methods for Analysis of Water 2016. Food Safety and Standards			
	Regulations Test Methods for Water/butter/cheese/milk	Authority of India (FSSAI), Ministry Of Health and Family Welfare			
	product for Processed Food Industry:	Government of India, New Delhi			
	(perform any two)				
	i. Proteolytic Plate Count				
	ii. Lipolytic Plate Count				
	iii. Thermophillic Bacterial Count (for Dairy Industry-Processing)				
	iv. Slime Forming Bacteria (for Dairy industry-Hot water				
4.	Food Safety and Standards Authority of India	1. Draft manual on method of microbiological testing (2016) microbiology of			
	(FSSAI)Regulations for Microbiological Testing of food:	foods. Food safety and Food Standards. Available			
	i. Detection and Confirmation of <i>Listeria monocytogenes</i> in Foods	at:https://old.fssai.gov.in/Portals/0/Pdf/Microbiological_Testing_Foods_Draf			
	ii. Fermentation Test (Incubation test for Cans, Tetrapacks, Standy	t_Manual_06_09_2016.pdf			

pouches).	2.	https://archive.fssai.gov.in/home/food-testing/food-testing-manual.html.
	3.	Manual for Good Food Laboratory Practices (GFLPs). 2018. Food Safety
		and Standards Authority of India (FSSAI), Ministry Of Health and Family
		Welfare Government of India, New Delhi

M.Sc. Microbiology Part II Semester IV

Guidelines for MBCP-4 Semester IV: Dissertation

- 1. A dissertation can be carried out by a single student or by group of students where the group should not contain more than two students.
- 2. The dissertation report will be prepared as per the thesis format.
- 3. Submission of the dissertation report will be at least ten days before the date of examination.
- 4. One copy of the report will be preserved in the department, in college.
- 5. If there are more than one student carrying out a single dissertation, a single report can be submitted to the department and these students will be assessed based on single oral presentation.
- 6. In such case, presentation should be carried out by all the students carrying out the same work; dividing the presentation equally among them.
- 7. At the time of presentation, the external and internal examiners appointed by the university will be present; the dissertation guide may or may not be present.
- 8. Presentation should be carried out to in the presence an audience comprising of examiners appointed by the university, departmental teaching staff and the postgraduate students of the department (M.Sc. I and II).
- 9. Oral presentation can be carried out using posters, blackboard, transparencies, model or LCD projector.
- 10. The allotted time for each oral presentation (one project) should be 10 to 12 minutes, followed by question and answer session of 5 to 8 minutes. The audience can participate in this session.
- 11. The assessment of the dissertation is for total of 100 marks (IA-30 and UA-70) out of which the university examinations assessment end semester will be for 70 marks and the in semester assessment will be for 30 marks.
- 12. The assessment of first 30 marks (in semester) will be carried out by the guide(s) who has supervised the work of the candidate(s) throughout the semester. The assessment will be carried out on the basis of the points, as per the accompanied format of the mark sheet. Head of the department should communicate this point wise assessment system to the dissertation supervisor, well in advance. Guide(s) will give appropriate marks, point-wise and

- submit it in a sealed envelope(s) to the Head of the respective department, three days prior to examination and project presentation. On the day of examination, Head of the department will hand over these unopened envelopes to the examiners.
- 13. Assessment of remaining 70 marks (end semester examination for both courses) will be carried out for individual student at the time of examination jointly by Internal and External examiners by the means of oral presentation. The assessment will be carried out on the basis of the points as per the accompanied format of the mark sheet.
- 14. Students should be made aware of the assessment parameters, on which they will be assessed throughout the semester and at the end of the fourth semester.
- 15. The external and internal examiners by mutual agreement will appropriately settle the marks given by the guide (reconsider, if necessary) and marks of oral presentation, and submit the mark lists to the Coordinator of the M. Sc. Examination Panel for that examination or directly to SPPU.

SAVITRIBAI PHULE PUNE UNIVERSITY

Practical Examination in M. Sc. Microbiology

Month Year

Course MB CP-4 (Dissertation)

Name of the center:
Name of the student:
Exam No.:
Point wise mark sheet – to be filled in by the Guide (Based on the evaluation carried out throughout the period of dissertation

Sr.	Points for Evaluation	Max.	Evaluation
No.		Marks	
1	Intellectual potential – Understanding of the research problem by the student (topic selection)	5	
2	Research aptitude –		
	a) Depth of literature survey for the proposed work.	3	
	b) Inputs of student in development of plans and protocols for the experimentation (methodology)	5	
	c) Ability to analyze data and formulate a solution (statistical analysis)	5	
	d) Analytical and reasoning abilities of the student for interpretation of data, inputs in discussion	5	
3	Motivation – punctuality, meeting dead-lines and seriousness (attendance)	2	
4	Ability to work with others	2	
5	Communication skill – oral and written (conferences, oral, ppt., publication)	3	
	Total	30	

Place of work	:

Name of the Guide :

Date and Signature

SAVITRIBAI PHULE PUNE UNIVERSITY

Practical Examination in M. Sc. Microbiology

Month Year

Course MB CP-4 (Dissertation)

Name of the center: _	 	
Name of the student:	 _	
Exam No.:		

Sr.	Points for Evaluation	Max.	Evaluation
No.		Marks	
1	Proficiency of presentation skills – use of audio-visual aids, preparation of graphs, charts, models, statistical analysis etc., use of scientific language	10	
2	Research potential of the work, results and interpretation, outcome of the study and possible future plans, publication potential of the work towards society	10	
3	The dissertation report preparation (scientific writing) and its contents	5	
4	Abilities of satisfactory responses to the queries from the audience (defense)	10	
	Total	35	

Point wise mark sheet – to be filled in by External examiner (Based on oral presentation and viva voce of the dissertation as end semester evaluation)

Place of work	:
Name of the External Examiner	:
Signature	:

Date

SAVITRIBAI PHULE PUNE UNIVERSITY

Practical Examination in M. Sc. Microbiology

Month Year

Course MB CP- 4(Dissertation)

Name of the center:
Name of the student:
Exam No.:
Point wise mark sheet – to be filled in by Internal Examiner (Based on oral presentation and <i>viva viva voce</i> of the dissertation as end semester evaluation)

Sr.	Points for Evaluation	Max.	Evaluation
No.		Marks	
1	Proficiency of presentation skills – use of audio-visual aids, preparation of graphs, charts, models, statistical analysis etc., use of scientific language	10	
2	Research potential of the work, results and interpretation, outcome of the study and possible future plans, publication potential of the work towards society	10	
3	The dissertation report preparation (scientific writing) and its contents	5	
4	Abilities of satisfactory responses to the queries from the audience	10	
	Total	35	

Place of work	:
Name of the Internal Examiner	:
Signature	:
Date	: