## Foundation for Clinical Practice (FCP)

## 25th June 2009

Topic & Concepts	Objectives	Time	Dept.	T/L activity	Comments
	At the end of the module, the student should be able,				
2006-3/SBM-1/01					
Introduction to Pathology	to understand the purpose of the module and the basis for the design of the module	1h	Patholo gy	Introductory session	
2006-3/SBM-1/02			37		
Introduction to Pharmacology -					
a. What is a drug?	define the following terms     Drug, Pharmacology, Therapeutics, Clinical Pharmacology,     Pharmacokinetics, Pharmacodynamics, "Medicines".	- 1h	1h Pharma cology	Lecture	Introduction to clinical pharmacology assignment
b. The need for the use of drugs in health care	compare and contrast 'drug' vs 'poison'     identify the broad principles of use of drugs in the management of common illness				and a state of the
2006-3/SBM-1/03	Common inness				A list of pre requiste knowledge for the FCP will be provided to the students.
Acute inflammation and suppuration	to define the process of acute inflammation.				
	2.to describe in detail* the various steps, controlling factors, sequale, complications and clinicopathological effects of acute inflammation. (includes suppuration)	9h Patholo	Patholo	Lectures (6h) +	
2006-3/SBM-1/04			gy	Museum Class (3h)	This will be supplemented by clinical lecture demonstration and will be done by the clinicians.
Chronic inflammation	1.to define the process of chronic inflammation     2. to describe in detail* the non-specific and specific types of chronic				detailed objectives are given separately to the students Application of wound healing in clinical practice by a
	inflammation, its sequele and complications				
2006-3/SBM-1/05					surgeon (1h) ** Pathology slide class - Review of Microscopic slides of
Wound healing	to describe the process of healing in injured tissue and its complications				general pathological processes
	2. to describe in detail* the process of healing in different types of tissue and surgical wounds.     3. to describe in detail* the formation of the organ of repair-namely	5h	Patholo gy	Lectures (4h) + Museum Class (1h)	
	granulation tissue.				
2006-3/SBM-1/06					This will be supplemented by clinical demonstrations in
Necrosis and apoptosis	to describe in detail* the morphological changes that occur in irreversibly injured cells and the clinico-pathological effects of such necrosis     to outline the non reversible types of cell injury.		4h Patholo gy		the wards during the introductory clinical appointments, and will be done by the clinicians. The clinicians will be informed of the topics during each week.  * detailed objectives are given separately to the students
	3. to describe in detail* the pathogenesis and pathology of different types of necrosis	4h		Lectures (3h) +	
	4. to outline the clinicopathological effects and recognition of necrosis			Museum class - (1h)	
	5. to define the term reperfusion injury and describe the process     6. to define the term apoptosis and discuss the clinicopathological significance				
	7. to name the steps in apoptosis and the controling factors  8. to differentiate apoptosis from necrosis				

2006-3/SBM-1/07					
Tuberculosis	1. to describe the pathogenesis of tuberculosis				
			Patholo		
	to understand the concepts of primary and postprimary tuberculosis	2h		Lectures (2h)	
	to describe the complications of the tuberculosis		gy		
	to explain pathological basis of the clinical effects				
2006-3/SBM-1/08					
Disorders of Growth and	to outline the ways in which different cell types react to increased work				
differentiation	demand and chronic irritation.			Lectures (3h) + Museum Class - (2h)	
	2. to describe in detail* the process of hypertrophy, hyperplasia, atrophy	5h	Patholo		
	and metaplasia, and the pathology of these processes.		gy		
	3. to give examples and also to state the clinico pathological effects of the				
2006-3/SBM-1/09	processes mentioned above.				
Thrombosis	to list the main factors which predispose to thrombosis.		1		
THIOHIDOSIS					
	2. to describe in detail* the pathogenesis and sequelae of thrombosis in		Patholo	Lectures (2h) +	
	different types of blood vessels and the heart, and the fate of thrombi	3h	gy	Museum class (1h)	
	3. to mention the clinicopathological features of thrombosis in the		97	Maccaill class (111)	
	different types of blood vessels.				
2006-3/SBM-1/10	71				
Embolism	1. to define the process of embolism.				
	2. to describe in detail* formation of different types of emboli and	6h	Patholo	o Lectures (2h) + Practical (4h)	Clinical lecture demonstration by a surgeon (1h)
	describe the outcome of the different types of embolism including the		gy		
	clinicopathological effects.				
2006-3/SBM-1/11					
Congestion, oedema and	to describe the effects in tissue, when the vascular blood supply alters,			Lectures (4h) + Museum class (3h)	including lymphoedema
infarction	and the venous return is hampered.				3,1
	2. to define the processes hyperaemia(active and passive), oedema and	7h	Patholo		
	infarction  3. to describe in detail* the pathogenesis of these processes.	. /11	gy		
	4. to list the processes that injure lymphatics and the clinicopathological				
	outcome due to injured lymphatics.				
2006-3/SBM-1/12	outcome due to injured lymphatics.				This will be supplemented by clinical demonstrations in
Amyloidosis	1. to define the process of amyloidosis.				the wards during the introductory clinical appointments,
, and the second	to describe in detail* the pathogenesis, types and clinical effects and	2h	Patholo gy	lo Lectures & Case discussions (2h)	and will be done by the clinicians. The clinicians will be
	methods of diagnosis of of amyloidosis.				informed of the topics during each week.
2006-3/SBM-1/13	,				* detailed objectives are given separately to the students
	1. to describe the process of pathological calcification and to state clinical		Patholo gy		
Other accumulations	examples.	2h		Lectures (2h)	
	2. to enumerate the types of abnormal pigments in the living persons and	211			
	their pathogenesis and clinical importance.				
2006-3/CLM-1/01					
Abnormal constituents in urine	to perform inward tests for urinary protein, sugar, bile and ketone bodies	4h		Bioche Practical (4h)	
	, and the second		mistry		
2006-3/SBM-1/14	1 to describe the right factors of atherescaleresis		1	1	
Atherosclerosis	1. to describe the risk factors of atherosclerosis     2 to describe the netheropoles and nethelogical processes involved in		Patholo gy		
	2 to describe the pathogenesis and pathological processes involved in atherosclerosis	2h		Lectures (2h)	
	3. to describe the complications and clinicopathological effects of	211		Lectures (2n)	
	atherosclerosis.				
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2006-3/SBM-1/15					
Neoplasia and Carcinogenesis					
a. Introduction to neoplasia and oncogenesis	to describe the fact that DNA alteration in a cell can lead to tumours and dysplasia.			Lectures (3h) + Museum class (1h)	This will be supplemented by clinical demonstrations and will be done by the clinicians (1h)
	2. to describe in detail* the process of carcinogenesis and concepts of dysplastic and premalignant lesions.     3. to describe in detail* the different types of tumours and their	4h	Patholo gy		
b. Spread of tumours	pathogenesis and morphology and differences in behaviour.  4. to describe in detail* the modes of spread of malignant tumours and the clinicopathological effects.	3h	Patholo	Lectures (2h) + Museum class (1h)	
2006-3/SBM-1/16			- 0,		
Antineoplastic Drugs					
	state how neoplastic cells/tissues differ from normal cells/tissues with respect to potential targets for drug therapy in neoplastic disease				
	2. explain the basis of				
	a. combination chemotherapy	2h	Pharma	Lecture	
	b. resistance to chemotherapy	211	cology	Lecture	
	c. adverse effects of chemotherapy				
	classify antineoplasitc drugs				
	4. describe the mechanism of action, pharmacokinetics, clinical uses,				
	adverse effects of commonly used antineoplastic drugs				
2006-3/SBM-1/17					
Applied general pathology	to define and explain the pathogenesis of erosions, ulcers, strictures and stenosis, blisters and bullae, fistula, sinus, polyps, adhesions, scars, fungating mass, organomegally, macule, papule, purpura, ecchymosis, naevi & warts and papillomata & application of these in systems	10h	Patholo gy	Lecture demonstration	clinical applications with clinicians
2006-3/SBM-1/18					
Drug action - Pharmacodynamics					Students are expected to work on computer assisted leaning package before and after a lecture
a. Modes of action of drugs at different levels :molecular, cellular, tissue/organ & overall individuals	list the mechanisms by which drugs exert chemical influences at cellular level to produce a pharmacological response				
	2. define and give examples of		Pharma cology/B iochemi stry		
	(I). receptor				
	(ii). drug binding sites				
	(iii). ligand	4h			
	(iv). agonist				
	(v). antagonist				
	(vi). partial agonist				
	(vii). inverse agonist				
	(viii), receptor affinity				
	(ix). receptor occupancy (x). spare receptors	<del> </del> 			
	(xi). efficacy				
	(xii). potency				
	(All). Policinoj		1		

b. Receptor as target for drug					
action	classify receptors based on their structure function	r			
	2. briefly explain the signalling mechanisms by which receptor activation	İ			
	is coupled to cellular effector systems.				
c. Targets for drug action	·	2h	Pharma	Lectures	
	(I). reversible/irreversible antagonism	211	cology	Lectures	
	(ii). competitive/non competitive antagonist				
	(iii). physiological antagonisms				
	(iv). tolerance, tachyphylaxis				
	(v). placebo and placebo effect				
2006-3/SBM-1/19					
Dose-response relationship:					
a. Drug dose-response     relationship-variations between     individuals	draw the concentration-effect curves for the relationship of the effect against				
	(I). (full) agonist concentration				
	(ii). logarithm of agonist concentration	3h	Pharma	Lectures	
	(iii). log-partial agonist concentration	ان ع	cology	Lectures	
	(iv). log full agonist concentration in the presence of a fixed dose+				
	increasing doses of competitive reversible antagonist				
	(v). log full agonist concentration in the presence of a competitive				
	irreversible antagonist				
	log full agonist concentration in the presence of a partial agonist				
b. Basis of adverse and toxic					
effects		01	Pharma		During hospital based assignment the studnts are
	1. define' adverse effects' of drugs	2h	cology	Lecture	expected to observe and record drug effects
	2. describe the mechanisms of adverse effects of drugs				
	explain how these reactions could be minimised/prevented				
	define therapeutic index	ł			
	<ol><li>describe the different mechanisms by which drugs may cause cell damage, cell death, mutagenesis, carcinogenicity and teratogenicity</li></ol>				
	list drugs that are potentially	1h	Pharma cology	Lecture	
	(I). hepatotoxic	İ			
	(ii). nephrotoxic				
	(iii). carcinogenic				
	(iv), teratogenic		<u> </u>		
c. Assessment & monitoring of	list the methods by which the effects of drug therapy could be				
drug effects	measured	2h	Pharma	Tutorials/ Lectures	During hospital based assignemtn the studnts are
	2. describe how the measurement of plasma drug concentrations helps		cology	. 3.33.5/ 20010100	expected to observe and record drug effects
	in monitoring drug therapy				
2006-3/SBM-1/20			1		
Pharmacokinetics			1		
How does the body handle					
drugs?		ł	Dhorns -		
a. Transport across cell	describe the mechanisms of transport of drug molecules across the		Pharma cology+		lipid/water solubility, diffusion, facilitated diffusion, active
I	cell membrane and the factors that influence such mechanisms.		Bioche		transport, efflux transporters such as ATP-binding
membrane:	ceii membrane and the factors that inhuence such mechanisms.		mistry		cassette (ABC) proteins, pinocytosis
		l	misuy		

b. Absorption: routes of administration	list different routes of administration of drugs				
	list the different types of dosage forms/special drug delivery systems     (eg. Metered Dose, Inhalar, Enteric coated formulation, spansules)				Assignment/skills lab activity
c. Distribution in tissues, body compartments and across barriers	describe the advantages and disadvantages of the routes mentioned in b.1 and drug dosage forms mentioned in b.2		5		
	list the different compartments of the body into which drugs are distributed		Pharma cology/B		
	describe the factors which influence the distribution of drugs into different compartments.		iochemi stry		
	explain the concept of redistribution of drugs	İ			
	explain the concept of barriers across tissues for transport of drugs				
d. Biotransformation	explain the basic mechanisms by which drugs undergo biotransformation in the body	6h+SGLA- CAL			Changes in different categories of population (elderly,pediatric and organ faliure)
	list the common drugs which induce/inhibit the cytochrome P 450	0,12		Lectures/Tutorials	
	enzyme system	ļ			
e. Elimination	1. define the terms "elimination" and "excretion"	<b>.</b>			
	<ul> <li>(I). state the physiological processes of different organ-systems that are involved in drug elimination</li> </ul>				
	(ii). explain the basic mechanisms by which drugs are excreted via				
f. Analytical pharmacokinetic	kidneys.  1. define the following	ł			
parameters	(I). bioavailability	1			
parameters	dosage regimen	1			
	(ii). bioequivalence	ł			
	(iii). first pass effect	ŧ	Pharma		
	(iv). Area Under the Concentrate-time curve (AUC)	t	cology/P		
	(v). (apparent) volume of distribution	•	hisiology		
	(vi). clearance	İ			
	(vii). half life	İ			
	(viii). steady state concentration	İ			
	(ix). loading dose	İ			
	(x). maintenance dose	İ			
	(xi). dosage regimen				
	explain the principles of calculating the bioavailability, volume of				
	distribution, clearance, loading dose & maintenance dose.				
g. First-order & Zero-order kinetics	explain first order kinetics and zero order kinetics	1h	Pharma cology	Lecture	
h. Drug concentration vs time					
curve in different dosing	draw the concentration-time curves for				
regimes		ļ			
	(I). single IV bolus injection				
	(ii). intermittent IV bolus injection	<b>.</b>			
	(iii). continuous IV infusion	2h	Pharma	Tutorials	
	(iv). single-dose oral administration	<b>ļ</b>	cology		
	(v). intermittent oral administration	<b>ļ</b>			
	(vi). modified-release formulations	1			
	(vi): modified release formulations	Ť			

2006-3/SBM-1/21					
Autonomic Nervous system					
	1. recall the anatomical and functional organisation of autonomic nervous system			Lecture/ Lutorial	
	recall the anatomy and the physiology of the cholinergic and the noradrenergic 'junctions.				
	3. recall the types of autonomic receptors with examples typical sites.				
	4. describe the mechanism of action, pharmacokinetics clinical effects of :				
	(I). cholinoceptor agonists	3h	Pharma		
	(ii). acetylcholinesterase inhibitors	,	cology		
	(iii). acetylcholinesterase re-activators				
	(iv). muscarinic receptor antagonists				
	(v). ganglion-blocking nicotinic antagonists				
	(vi). adrenoceptor activators				
	(vii). adrenoceptor antagonists				
	5. describe the important clinical uses, adverse effects, toxic effects,				
	contra-indications with regard to the drugs acting on the autonomic				
2006-3/SBM-1/22	nervous system.				
Pain Control					
a. Physiology of pain					
a. r nyolology or pani	recall the definition of pain and briefly explain theories of pain				
	2. classify pain				
	recall physiology of pain perception (Stimuli, receptors, pathways and				
	central connection)				
	4. list methods of pain relief	•			
	5. classify pharmacological agents used in pain relief (with main indications)				
	6. explain the basis of neuropathic pain.	•	Pharma		
b. Opioid Analgesics		•	cology	Lecture	
	classify the agents acting on opioid receptors	•	0,		
	describe the mechanism of action of opioid analgesics.	01			
	describe the pharmacokinetics of the drugs acting on opioid receptors	2h			
	describe the adverse effects of opioid analgesics.				
	list the clinical uses of opioid receptor antagonists				
c. Non-steroidal anti- inflammatory drugs (NSAIDs)					
	describe the physiological/pathological roles of Cyclo-oxygenase-1     (COX - 1) and COX - 2 enzymes.				
	describe the pharmacokinetics, clinical uses, important adverse effects and drug interactions of NSAIDs (including COX - 2 inhibitors).				
	3. list the commonly used NSAIDs.	•			
2006-3/SBM-1/23	,				
Introduction to Haematology	recall the cellular components in blood and haemopoiesis	6h	Patholo gy	Lectures (6h)	
	outline the common types of non malignant and malignant diseases of blood	011			
2006-3/SBM-1/24					
Introduction to Clinical Pathology	outline the applications of serological and haematological investigations in patient management	1h	Patholo gy	Lecture (1h)	