





#### **COURSE SPECIFICATION**

### (Advanced Medical Biochemistry & Molecular Biology)

#### (Basic level II)

#### Faculty of Medicine- Mansoura University

#### (A) Administrative information

(1) Programme offering the course:	Master degree of Basic Medical Sciences in Biochemistry
(2) Department offering the programme:	Medical Biochemistry Department
(3) Department responsible for teaching the course:	Medical Biochemistry Department
(4) Part of the programme:	Second part
(5) Date of approval by the Department's council	29/4/2018
(6) Date of last approval of programme specification by Faculty council	S to
(7) Course title:	Advanced Medical Biochemistry & Molecular Biology (Basic level II)
(8) Course code:	BIC 504ad
(9) Total teaching hours:	Lectures:195 hrs Practical: 240 hrs
(10) Total credit hours:	13 hours Lectures 8 hours practical

## (B) <u>Professional information</u>

#### (1) Course Aims:

Provide the candidate with a recent knowledge in modern biochemistry and molecular biology, enabling recent graduates from the Master's program to found technical positions in academic and labs, pursued careers in teaching, science writing and editing, or have gone on to advanced studies for the PhD. Degree.

(2) Intended Learning Outcomes (ILOs): On successful completion of the course, the candidate will be able to:

A-	Knowledge and Understanding:
All.1	All.1.1 Define water as an Ideal Biologic Solvent
	All.1.2 Study Covalent & Noncovalent Bonds Stabilize Biologic Molecules
	All.1.3 Define PH &its calculation
	All.1.4 Describe buffer system in the body&its function
All.2	All.2.1 define classification and strucrure of monosaccharides
	All.2.2 define derivatives of monosaccharides
	All.2.3 define disaccharides
	<b>All.2.4</b> define polysaccharides( classification, homopolysaccharides, heteropolysaccharides "GAGs"
	All.2.5 Describe strucure, importance and classification of lipids
	<b>All.2.6</b> Discuss structure and function of phospholipids, glycolipids, sulpholipids and lipoproteins
	All.2.7 Define structure and function of derived lipids
	All.2.8 recognize chemistry and function of carotenes
	All.2.9 Describe importance, classification, general properties of protein
	All.2.10 Describe Structure, classification, properties of amino acids
	All.2.11 Describe protein strucure, protein folding and protein misfolding
All.3	All.3.1 Describe digestion, absorption of CHO and glucose uptake by tissues.
	<b>All.3</b> .2 Describe glycolysis (definition, site, steps, biomedical and clinical importance, regulation, energetic and clinical aspects)
	All.3.3 discuss gluconeogenesis (defination, site, importance, gluconeogenic substrates, steps, regulation)
	<ul><li>All.3.4 discuss glycogen metabolism</li><li>Structure and function of glycogen</li></ul>
	Glycogenesis (definition, site and steps)
	Glycogenolysis (definition, site and steps)
	Regulation of glycogen metabolism
	Glycogen storage disease
	AII.3.5 Recognize Pyruvate metabolism with stress on Oxiadative decarboxylation (definition, site, steps, regulation)

	All.3.6 Describe citric acid cycle(definition, site, steps, biomedical importance, regulation and inhibitors, energetic, clinical aspects and role of vitamins)
	<b>A</b> II.3.7 discuss hexose monophosphate shunt (definition, site, biomedical importance, function of NADP, regulation and clinical aspects)
	All.3.8 discuss uronic acid pathway(definition, site, importance, pathways of UDPG, biosynthesis of amino sugars)
	AII.3.9 Discuss Metabolism of mono and disaccharides
	<ul> <li>Fructose metabolism (biomedical importance, conversion of fructose to glucose, conversion of glucose and mannose to fructose and inborn errors of fructose metabolism)</li> </ul>
	<ul> <li>Galactose metabolism (biomedical importance, conversion of galactose to glucose, conversion of glucose to galactose and inborn errors of galactose metabolism)</li> </ul>
	All.3.10 Describe Insulin (Structure, synthesis, mechanism of action, regulation of secretion, metabolic effects and catabolism)
	All.3.11Describe Glucagon (Structure, mechanism of action, regulation of secretion and metabolic effects)
	All.3.12 Interpret Blood glucose level (regulation of blood glucose level and clinical aspects; glucosurira, hyper and hypoglycemia)
	All.3.13 define Glucose homeostasis (Regulation of blood glucose) Glycoproteins (structure, synthesis, degradation and biomedical and clinical
	importance)
All.4	
	All.4.1 Describe and differentiate between types of Diabetes milletus
	<b>All.4.2</b> Define diabetes mellitus and know its incidence, pathogenesis, metabolic changes, diagnosisand treatment of both types
	All.4.3Explain Complications of diabetes mellitus (acute and chronic and its pathogenesis)
All.5	All.5.1Describe Lipogenesis (definition, site, regulation, steps)
	All.5.2Discuss Fatty acid synthsis
	<ul> <li>Synthesis of saturated FA (Cytoplasmic FA synthesis, Mitochondrial FA synthesis, Microsomal FA synthesis)</li> </ul>
	Synthesis of unsaturated FA
	Synthesis of glycerol and TG
	AII.5.3Describe and compare between different types of Fatty acid oxidation

	<ul> <li>B_oxidation, Alpha oxidation, Omega oxidation (definition, site, steps)</li> </ul>
	Oxidation of unsaturated FA
	All.5.4 Recognize Active acetate (sources and fate)
	All.5.5 Discuss Ketone bodies metabolism
	Ketogenesis (definition, site, steps, biomedical importance and regulation)
	Ketolysis (definition, site, steps, biomedical importance and regulation)
	Ketosis (definition, pathogenesis, causes and effects)
	All.5.6 Describe Lipoprotein metabolism and differentiate between different types of lipoproteins
	<ul> <li>Definition, site, steps, biomedical importance, regulation and metabolism of each type</li> </ul>
	<ul> <li>Apoproteins ( definition, role and types)</li> </ul>
	Enzymes in lipid transport
	Primary disorders of plasma lipoproteins.
	All.5.7 Discuss Eicosanoids metabolism (Definition, members, synthesis, biological actions, clinical aspects)
	All.5.8 Describe Cholesterol metabolism
	<ul> <li>Structure, Synthesis, Transport and Degradation</li> </ul>
	<ul> <li>Blood cholesterol levels and its clinical aspects</li> </ul>
	<ul> <li>Bile acids and bile salts (structure, synthesis and clinical aspects)</li> </ul>
	<ul> <li>Steroid hormones (synthesis, secretion and mechanism of action)</li> </ul>
	All.5.9 Recognize Phospholipid and glycosphingolipids metabolism
	Structure, Function, Biosynthesis and catabolism of different types of PL
	Types and synthesis of glycosphingolipidos
	Sphingolipidosis
	All.5.10 Describe role of adipose tissue in lipid metabolism with stress on hormonal regulation
	All.5.11 Discuss Fatty liver (definition, causes ,pathogenesis and lipotropic factors)
AII.6	All.6-1 describe amino acid pool
	All.6-2 demonstrate catabolic pathways of amino acids (transamination-deamination-decarboxylation-transamidation)
	All.6-3 Describe sources & fates of ammonia

	All.6-4 Describe urea biosynthesis (steps-regulation-metabolic disorders)
	All.6-5 Describe the nitrogen balance
	All.6-6 recognize biosynthesis of non essential amino acids
	All.6-7 discribe catabolism of carbon skeleton of a.a.
	All.6-8 Describe conversion of amino acids to specialized product
	All.6-9 list nitrogen containg compound.
All.7	All.7-1 Describe synthesis of purine nucleotide
	a-denovopathway (steps-regulation)
	b-salvage pathway
	c-deoxyribonucleotide synthesis(steps-regulation)
	All.7-2 discribe catabolism of purin nucleotides
	All.7-3 explain metabolic disorders of purine metabolism(hypouricemia- hyperuricemia)
	All.7-4 Describe pyrimidine synthesis & degradation
	a-denovopathway (steps-regulation)
	b-salvage pathway
	c-catabolism
	All.7-5 synthetics base analoges used in chemotherapy
All.8	All.8-1 describe (structure-synthesis-regulation-disorders) of porphyrin
All.9	All.9-1 Describe enzyme change in fed & fasting state
	All.9-2 discribe role of (liver-adipose tissue-muscle-brain)in fed &fasting state
All.1	All.10.1 describe structure & funtion of plasma membrane
U	All.10.2 Describe Artificial membrane
	Transport processes
	Transport proteins
	All.10.4 enumerate membrane Diseases
All.1	Steroid synthesis
	Catecholamines synthesis
	<ul> <li>Thyroid synthesis</li> <li>Hormono synthesis from larger peptide presureers</li> </ul>
All.1	Hormone synthesis normarger peptide precursors
2	AILTZ.T Describe The Target Cell Concept

	All.12.2 Discuss Hormone Receptors(Definition/ nature/ characters/types)
	All.12.3 Describe Classification of hormones: according to:
	Chemical nature
	Mechanism of Hormonal action
All.1 3	All.13.1 enumerate immunoglobulins types, Functions, structure& Diseases
Ū	All.13.2 describe collagen Structure, types, biosynthesis& degradation
	All.13.3 explain elastin Structure, degradation & disorders of degradation
	All.13.4 understand Fibrillin& Marfan Syndrome
	All.13.5 discuss Fibronectin structure& Functions
AII.1 4	<b>All.14</b> .1 Describe fat- soluble vitamins ( active forms , synthesis ,function ,toxicity , deficiency ) .
	All.14.2 Describe water –soluble vitamins (chemistry ,function ,toxicity , deficiency & co-enzyme forms of each one of vitamin B complex members ).
	All.14.3 Describe absorption ,transport , storage , function ,toxicity & deficiency of macroelements & trace elements
All.1	All.15.1 Discuss mechanism of action of enzymes.
5	All.15.1a to identify characters and types of enzymes.
	All.15.1b to compare different mechanisms to facilitate catalysis.
	All.15.2 Identify role of prosthetic groups, cofactors and co-enzymes and their types.
	All.15.3 identify isoenzymes
	All.15.4 explain how catalytic activity of enzymes facilitate their detection.
	All.15.5 demonstrate application of enzymes in diagnosis of diseases.
	All.15.6 illustrate role of recombinant DNA in studying enzymes.
	All.15.7 describe types of chemical reactions.& factors affecting the reaction rate
	All.15.8 recognize kinetics of enzymatic catalysis.
	All.15.8 a. Factors affecting the rate of enzyme-catalyzed reactions.
	<b>All.15.8</b> b. To demonstrate effects of substrate concentration by Michaelis- Menten and Hill equation.
	All.15.8 cTo distinguish competitive and non competitive inhibition.
	All.15.9 define types of enzyme-catalyzed reactions.
	All.15.10 recognize role of enzymes in drug discovery.
	All.15. 11 Describe How to explain regulation of enzyme at both quantity & catalytic activity levels .

All.1 6	All.16.1 Describe Important Proteins of Muscle
•	All.16.2 Explain the sequence of Events in Contraction and Relaxation of Skeletal Muscle & Smooth Muscle
	All.16.3 Discuss Regulation of Muscle Contraction
	All.16.4 Describe Role of Ca2+ in Regulation of Muscle Contraction
	All.16.5 Describe Role of Sarcoplasmic Reticulum in Regulation of Intracellular Levels of Ca2+ in Skeletal Muscle
	All.16.6 describe Channelopathies& Inherited Cardiomyopathies
	All.16.7 Describe synthesis& Functions of Nitric Oxide
	All.16.8 enumerate Mechanisms Replenish Stores of ATP in Muscle
	<b>All.16.9</b> Describe Types of Muscle Fibers and Major Fuel Sources Used by a Sprinter and by a Marathon Runner
	All.16.10 study of Cytoskeleton (Intermediate Filaments, Microfilaments & Microtubules).
All.1 7	<b>All.17</b> .1 Describe electron transport chain( def, components, enzyme complexes, sequence of events)
	<b>All.17</b> .2 Describe oxidative phosphorylation (Definition, site , energetic, Theories, p/o ratio,
	Inhibitors& Uncouplers)
	All.17.3 Differentiate between energy production in biological and non biological systems
	All.17.4 Mention Genetic mitochondrial disorders
	All.17.5 discuss oxidation of cytoplasmic NADH
	<b>All.17</b> .6 Discuss and interpret bioenergetics (definition, first law of thermodynamics, gibbs free energy and standard free energy)
	All.17.7 Recognize ATP (sources and biological importance)
	All.17.8 Describe Low and high energy bonds
	All.17.9 Describe the redox potential.
	All.17.10 Discuss oxidoreductases
AII.1 8	A34.1Describe the two phases of Xenobiotics Metabolism
•	A34.2 Discuss Isoforms of Cytochrome p450&their function
	A34.3 Enumerate Responses to Xenobiotics
All.1 9	All.19.1Describe sources& effects of free radicals

	All.19.2 Explain protection against free radicals by Antioxidants
All.2	All.20.1 study Plasma Proteins & their Functions
0	All.20.2 discuss Iron metabolism (absorbtion and storage)
	All.20.3 explain Copper metabolism (absorbtion and storage)
	All.20.4 discuss the deficiency of alpha1-Antitrypsin as one of Plasma Proteins
	All.20.5 Describe Amyloidosis
	All.20.6 describe Hemostasis & Thrombosis (Phases, Types of Thrombi& Intrinsic &
	Extrinsic Pathways)
	All.20.7 Describe the Functions of the Proteins Involved in Blood Coagulation.
	All.20.8 enumerate Hereditary Bleeding Disorders
	All.20.9 describe Activation of Platelets & role of Aspirin as an Antiplatelet Drug
	All.20.10 discuss the regulation of Circulating Thrombin with stress on
	Antithrombin& Coumarin Anticoagulants
	All.20.11 describe role of Plasmin in Hemostasis
	All.20.12 explain the role of Endothelial Cells Hemostasis
	All.20.13 enumerate Laboratory Tests Measure Coagulation, Thrombolysis, &
	Platelet Aggregation
All.2	All-21.1 Discuss different levels of DNA structure (DNA primary&
1	different types of DNA secondary structure).
	All-21.2 Recognizing the definition & mechanisms of DNA denaturation &
	renaturation as methods of analyzing DNA structure.
	All-21.3 Differentiation between DNA & RNA.
	All-21.4 Enumerat different types of RNA with explaining the structure&
	function of each one.
	All-21.5 Discuss the chromatin structure with explanation of the histone
	proteins, nucleosomal structure & chromatin structure.
	All-21.6 Describ the high order structure of the chromatin & its incorporation
	in the activity state of the chromatin.
	All-21.7 Explain the DNA repetitive & non repetitive regions.
	All-21.8 Compare different types of rearranging the genetic material
	transposition with explain examples on each in the living system)
	All-21 0 Discuss DNA replication
	All-21.9 Defining the replication with recognizing the steps of replication
	<b>All-21.9a</b> Defining the representation with recognizing the steps of representation. <b>All-21.9b</b> Differentiating types of DNA polymerase ineukaryotes & prokaryotes &
	the function of each one
	<b>All-21.9c</b> Recognizing the replication polarity.
	<b>All-21.9d</b> identifying Timing of replication in the cell cycle & its application.
	All-21.10 discuss DNA mutation & repair.
	All-21.10a Recognizing the definition, causes & effect of DNA mutation.
	All-21.10b mention different types of DNA repair.
	All-21.10c Explain the clinical conditions associated with impaired repair.
	All-21.11: Discuss RNA synthesis:-
	All-21.11a Describe the classification of RNA and how it is synthesized from
	DNA template by RNA polymerase.

**All-21.11b** Compare between eukaryotic and prokaryotic DNA dependent RNA polymerase (also know the difference between euik and prok promoters).

**All-21-11c** Explain how RNA synthesis is cyclical process involve RNA chain initiation, elongation and termination.

**All-21.11d** Discuss different transcription factors and discusse the components, formation, assembly of basal transcription complex.

**All-21.11e** Recognize signals that regulate transcription termination and clarify how termination occurs either by Rho factor independent or Rho factor dependent manner.

All-21.12: Mention how RNA molecules usually processed before they become functional :

**All-21.12a** Discuss how introns removed and exons are spliced together and explain mechanism of mRNA splicing, alternative splicing and alternative promoters provides a form of regulation of mRNA.

All-21.12b Recognize how tRNA, rRNA are processed.

All-21.13 discuss how RNA is modified after it's synthesis.

All-21.13a Recognize copping, tailing, splicing and RNA editing of mRNA.

All-21.13b discuss post transcriptional modification of tRNA and rRNA.

All-21.13c Explain how RNA can acts as a catalyst.

**All-21.13d** discuss micro-RNA synthesis & their role in quality control of mRNA in P bodies.

**All.21.14** Recognize molecular basis of protein synthesis and how to determine the expression of certain proteomics in diseased cell:

All.21.14.a : Describe different features of genetic code

**All.21**.14.b: Explain the three phases of protein synthesis: initiation with formation of initiation complex, elongation and termination.

All.21.14.c : Understand regulation and control of protein synthesis

- Explain control at the level of gene expression.
  - Know regulation and control of initiation
  - Understand how protein synthesis respond to environmental threats.
  - Explain how viruses can affect protein synth.
  - Recognize post translational processing affects the activity of synthesized protein.
  - Describe the effect of Antibiotics on bacterial protein synth.

All.21.15 know updates of the regulation of gene expression

All.21.15.a Classifying types of genes according to the mechanism of their expression.

All.21.15.b recognizing different types of regulation of gene expression in

prokaryotes	
<ul> <li>Recognizing catabolic regulation (lac operon)</li> </ul>	
Explain co-repression (tryptophan operon)	
• Demonstrate genetic switching ( $\lambda$ phage cycle)	
All.21.15.c classifying the levels of euokaryotic regulation of gene expression.	
•regulation of gene expression at the Genomic level (gene rearrangement & gene amplification)	
•regulation of gene expression at the transcriptional level (DNA	
regulatory Protein& DNA regulatory regions) with explaining silence	r,
enhancer, Locus control region & insulator as DNA regulator regions	S.
<ul> <li>regulation of gene expression at Post transcriptional level (RI processing, RNA stability, RNA editing &amp; the effect of micro-RNA mRNA)</li> </ul>	NA on
Describe the basis of determining genes and methods of amplification	of
those genes:	
All.21.16 Describe Recombinant DNA technology	
<ul> <li>Explain clearly what is cloning and its steps</li> </ul>	
<ul> <li>Understand practical applications of recombinant DNA tech. a appreciate how molecular biology gives us new perspectives a</li> </ul>	and
<ul> <li>Differentiate between in vivo &amp; in vitro amplification(PCR) a explain applications of PCR.</li> </ul>	s and
All.21.17 Describe genomic technologies :	
<ul> <li>Compare between genomic and cDNA libraries.</li> </ul>	
<ul> <li>Understand different methods gene localization and generation</li> </ul>	ene
sequencing.	
<ul> <li>Know RNA and protein proming and protein – DNA interact mapping</li> </ul>	ion
All.21.18 Explain the role of gene therapy as a therapeutic indications of D	NA
technology :	
<ul> <li>Discuss types of diseases can be treated with gene therapy a</li> </ul>	and
types of vectors used.	
Understand well gene therapy strategies	
AII.2 AII.22.1 Define oncogenes&their role in cancer development	
All.22.2 List tumor markers&their use in dignosis andfollow up of cancer	
All.2 All.23.1 Understand the concept of stem cells and their importance.	
All.23.2 Summarize the causes of the major disorders affecting red blood cells.	
All.23.3 Discuss the general structure of the red blood cell membrane.	
All.23.4 Know the biochemical bases of the ABO blood group substances.	

	All.23.5 Indicate the major biochemical features of neutrophils and understand the
	basis of chronic granulomatous disease.
	All.23.6 Appreciate the importance of integrins in health and disease.
AII.2 4	<ul> <li>All-24.1. Discuss intraceller trafficking</li> <li>All-24.1.a. Identify the meaning of trafficking with stress on final protein distenations.</li> <li>All-24.1.b. Discuss the sequences of molecules carried by proteins to target them to specific organelles.</li> <li>All-24.1.c. Discuss the techniques used to study intracellular trafficking.</li> <li>All-24.2. Discuss Protien sorting</li> <li>All-24.2.a. Mention the different sorting branches (according to the site of protein synthesis).</li> <li>All-24.2.b Discuss cytosolic sorting branch (Mitochondrial, Nuclear &amp; Peroxisomal).</li> <li>All-24.2.c Discuss R.E.R sorting branch (Discuss the signal hypothesis, Mention several routes by which proteins are inserted into the membrane of E.R., Explain the role of G.A in retrograde transport )</li> <li>All-24.2.c Discuss Unfolded protein response (URP) &amp; E.R degeneration (ERAD) of protein.</li> <li>All-24.2.g Define the transport vesicles with special stress on its role on intracellular protein trafficking.</li> <li>All-24.2.h Explain the process of membrane assembly.</li> <li>All-24.2.h Explain the process of conformational diseases.</li> </ul>
All.2 5	All.25.1 Discuss apoptosis All-25.1.a. Definition- mechanism- importance of apoptosis.
	All-25.1.b. Discuss role of apoptosis in endoplasmic reticulum homeostasis.
	All-25.1.c. Discuss abnormalities of apoptosis that occur in disease states.
-	B-Intellectual skills:
BII. 1	Interpret symptoms, signs and biochemical laboratory findings of some metabolic disorders.
BII. 2	Point-out the etiology of metabolic disturbance in a given case study report.
BII. 3	Point-out the application of molecular biology in basic and clinical sciences.
BII. 4	Interpret symptoms, signs & biochemical laboratory findings of vitamins deficiency diseases
BII. 5	Interpret the clinical significance of determination of plasma levels of glucose, total proteins, SGOT, SGPT, bilirubin, albumin, cholesterol, TG, creatinine and uric acid
BII. 6	Diagnose the type of abnormality of pathological glucose tolerance curve

	B- Professional/practical skills:	
CII. 1	Perform DNA extraction by spin column.	
CII. 2	Perform molecular biology techniques: electrophoresis and conventional PCR.	
CII. 3	Identify gene polymorphisms by restriction endonucleases.	
CII. 4	Measure some parameters by ELISA.	

#### **D-** Communication & Transferable skills:

DII.1	To be able to work effectively in a group in lab or during preparation of seminars
DII.2	To respect the role of staff and co-staff members regardless of degree or
	occupation.
DII.3	To be able to use computer and IT.

# (3) Course content: Total teaching hours: 195 hr (13 credit hr), practical 240 hr (8 credit hr) A. lectures:

Subject	NO. of hours					
Module 1						
1.Water metabolism & acid-base balance	10					
2. Chemistry of carbohydrate, lipid & protein	20					
3.Metabolism of Carbohydrate& glycoprotein.	20					
4. Diabetes mellitus	5					
5 Metabolism of lipid & ecosanoids	20					
<b>6.</b> . Metabolismof protein & individual amino acids.	15					
7. Purine & pyrimidine nucleotides metabolism	7					
8. Porphyrine metabolism & bile pigment.	7					
<b>9.</b> Metabolic integration& the Fed/Fast cycle.	3					
10. Biological Transport and cell membrane	5					
<b>11</b> . Biochemistry of endocrine glands.	10					
<b>12.</b> Hormonal action & Cell signaling.	3					

Module 2		
1. Molecular biology & biotechnology	22	
2. Oncology (oncogenes & tumor markers).	4	
3. Immunochemistry and tissue chemistry	2	
4. Micronutrients (Vitamins & Minerals)	10	
5. Enzymes	10	
6. Muscle chemistry & metabolism	4	
7. Biological oxidation & bioenergetics.	5	
8. Xenobiotics and detoxification	2	
9. Free radicals and antioxidants	2	
<b>10.</b> Blood (plasma proteins & homeostasis).	4	
<b>11.</b> Protein traffic and sorting	4	
12. Apoptosis	1	
Total teaching hours	195	

#### B. Log book activities:

Subjects	NO. of hours
A. practical:	
1 DNA extraction (spin columns).	45
2 Electrophoresis (vertival and horizontal).	50
<b>3</b> Molecular biology techniques: conventional PCR.	50
4 Gene polymorphisms by restriction endonucleases.	45
<b>5</b> Measurement of some parameters by ELISA.	50

#### (4) Teaching methods:

- 4.1: Lecture
- 4.2: Practical class
- 4.3: Small group discussion with case study and problem solving
- 4.4: Tutorial
- 4.5: Seminars
- 4.6: Workshops

#### (4) Assessment methods:

5.1: MCQ Examination for assessment of knowledge and intellectual ILOs

5.2: Written Examination for assessment of knowledge and intellectual ILOs

**5.3: Oral examination for assessment of** ILOs number: knowledge and intellectual and transferable ILOs.

**5.4: OSPE Practical examination for assessment of** knowledge, intellectual, practical and transferable ILOs.

**5.5: Log book for activities for assessment of** : mainly for assessment of practical & transferrable skills which are accepted through attending different conferences, thesis discussions, seminars, workshops, attending scientific lectures as well as self learning.

**5.6:** The supervisor require certain lab tests or exam that are evaluated and signed by the supervisors in the log book (without marks).

**5.7: seminars:** the candidate should prepare and present at least one seminar in a topic related to the course and determined by the supervisors in front of the department staff (without marks).

#### Assessment schedule:

• After 36 month from job registration (written, oral and practical exam with marks).

#### Percentage of each Assessment to the total mark:

Tools	Marks	Percenta	ge of the	total mark	
Advanced Medical E	Biochemistry &	Molecular	Biology	(Basic level	
Written exam	300	46.3%			
Oral exam	150	23%			
Practical exam	150	23%			

#### Other types of assessment

• Log book required activities to go through 2<sup>nd</sup> part examination .

#### Other assessment without marks:

• Practical tests and/or exam as well as the seminarthroughout the course and lab rotation (without marks).The candidate should prepare and present at least one seminar in atopic related to the course and determined by the supervisors in front of the department staff (without marks)

#### (5) References of the course:

#### 6.1: Hand books:

• Medical biochemistry department (student book)

#### 6.2: Text books:

- Harper's Illustrated Biochemistry: 28th edition by Murray RK, Granner DK, Mayes PA, Rodwell VW, McGraw-Hill companies New York, 2009.
- Lippincott's Reviews of Biochemistry, 4th edition by Champe PC, Harvey RA, Ferrier DR, Lippincott William & Wilkins London, 2008.
- Textbooks of Medical Biochemistry, 7th edition by Chatterjea MN. and Shinde R. JAYPEE BROTHERS. New Delhi, India, 2007.
- Text book of Biochemistry with Clinical Correlations 5th Ed, Devlin TM Ed.Wiley -Liss New York 2002
- Zilva Clinical Chemistry & Metabolic Medicine, 7th edition, Crook MA, Hodder Arnold, London, 2006
- Pretest Biochemistry and Genetics: 3<sup>rd</sup> edition by Golder N. Wilson,library of congress cataloging-in-publication data, Dallas, Texas, 2007.
- Multiple Choice Questions in Biochemistry :2<sup>nd</sup> edition, by RC Gupta, JAYPEE BROTHERS. New Delhi, India, 2004.
- Case Files Biochemistry :2<sup>nd</sup> edition, by TOY SEIFERT STROBEL HARMS, McGraw Hill, USA, 2008.
- Board Review Series :Biochemistry, Molecular Biology and Genetics :5<sup>th</sup> edition, by T.A.Swanson, S.I.Kim, M.J.Glucksman, M.A.Lieberman, Lippincott William & Wilkins, Baltimore, USA, 2010.
- 6.3: Websites:http://www.medlib.iupui.edu/ref/biochem.htm
- The Biology Project (from the University of Arizona): <u>http://www.biology.arizona.edu/default.html</u>
- Harvard Department of Molecular & Cellular Biology Links: <u>http://mcb.harvard.edu/BioLinks.html</u>

#### (7) Facilities and resources mandatory for course completion:

- Lecture rooms: available in the department
- Laboratories: The Department has 3 laboratories for research with a wide range of instrumentation that is available for training and research .
  - library
  - Computer laboratories with a wide range of software
  - Intranet with a wide range of learning support material

Course coordinator: staff members of the credit

## Head of the department: Prof. / Ayman El-Baz

Date: 29/4/2018