

Degree programme	Type	Course
Microbiology	OP	4

Contact lecturer

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Group languages

You can consult this information at the [end](#) of the document.

Prerequisites

The prerequisites to follow the course are a working knowledge of the material of the courses "Biochemistry", "Animal Biology" and "Instrumental Techniques", and have extensive knowledge of Human Physiology and Human Metabolism.

Objectives

Clinical Biochemistry is an optional course of the 4th year of the Degree in Microbiology that is part of the Degree in Biomedical Sciences.

Clinical Biochemistry aims to initiate the student in understanding the *in vitro* study of the biological properties that contribute to the prevention, diagnosis, prognosis and monitoring of diseases and disease states in humans.

The general objectives of the course are:

- 1) Provide students with knowledge of the specific characteristics of a clinical biochemistry laboratory.
- 2) Describe the pathophysiology and molecular basis of the most prevalent diseases.
- 3) Identify the analytical methods commonly used in the clinical laboratory.
- 4) Describe how can contribute the clinical laboratory to assess the health status of individuals.

At the end of the course the student will know: the pathophysiological bases of the most relevant and prevalent diseases in our population; the main biological properties that are altered in these diseases and are examined in a clinical biochemistry laboratory; the procedures for the biological properties measurement and test; and their semiologic characteristics. It will also be familiar with the use of tools for the operation in a clinical biochemistry laboratory: instructions or work protocols, implementation of internal control quality program, participation in an external quality evaluation programs and use of automated measurement systems.

Learning outcomes

- CM19 (Propose methods and procedures within the field of biochemistry, physiology and biotechnology to provide innovative responses to the needs and demands of society, and valuing their social, economic and environmental impact.) Propose methods and procedures within the field of biochemistry, physiology and biotechnology to provide innovative responses to the needs and demands of society, and valuing their social, economic and environmental impact.
- CM20 (Integrate knowledge of biology and biochemistry to develop an academic and professional work, and its presentation in writing or orally and publicly, working individually and in teams.) Integrate knowledge of biology and biochemistry to develop an academic and professional work, and its presentation in writing or orally and publicly, working individually and in teams.
- KM29 (Identify the physiopathological bases of the most relevant diseases with the highest prevalence in our population.) Identify the physiopathological bases of the most relevant diseases with the highest prevalence in our population.
- SM29 (Interpret biochemical and physiological parameters used for screening, diagnosis, prognosis or monitoring of different pathologies or pharmacological studies.) Interpret biochemical and physiological parameters used for screening, diagnosis, prognosis or monitoring of different pathologies or pharmacological studies.

Contents

Theme I. Metrological and semiological aspects of clinical laboratory

INTRODUCTION. Concepts. Pre-analytical, analytical and post-analytical phases of the clinical laboratory. Collection, preparation and preservation of specimens. Pre-analytical variability.

METROLOGY. Values and quantities, observations and measurements. Measures and errors. Random error: precision. Systematic error: trueness. The true value. Calibration and traceability. Specificity. Error of measurement: accuracy and uncertainty. Measuring analytical range. Analytical sensitivity. Detectability.

QUALITOLOGY. Control materials. Basis of internal control: control rules and charts. Algorithms. External quality assessment programs.

BIOLOGICAL VARIABILITY AND REFERENCE VALUES. Intra- and inter-individual biological variability. Theory of reference values. Transversal and longitudinal comparisons. Production of reference values.

SEMIOLOGY. Discriminant capacity. Cut-off value. Diagnostic sensitivity and specificity. Prevalence and predictive value. Likelihood ratio. Receiver operating curves (ROC).

Theme II. Biochemical assessment of metabolic pathways

PROTEINS. Classification of plasma proteins. Methods of identification, detection and quantification. Identification of electrophoretic profiles. Hyperproteinemia and hypoproteinemia. Polyclonal and monoclonal hyperimmunoglobulinemia. Paraprotein.

ENZYMES. Diagnostic usefulness of measuring plasma enzymes. Measuring the catalytic concentration and the mass concentration. Transformation and factors that affect it. Calibration. Standardization measures. Major diagnostic enzymes of interest.

CARBOHYDRATES. Hormonal control of glucose homeostasis. Hyperglycemia and diabetes mellitus and

alterations in glucose tolerance. Procedures for measuring glucose, hemoglobin A1c and albumin in urine.

LIPOPROTEINS. Structure and classification of plasma lipoproteins. Procedures for the study of dyslipidemia: total cholesterol and triglyceride. Separation of lipoproteins, VLDL cholesterol, HDL and LDL. Apolipoproteins. Classification of dyslipidemia. Genetic basis. Atherothrombosis: coronary heart disease and risk factors.

CALCIUM. Hormonal regulation of calcium homeostasis. Hypercalcemia and hypocalcemia. Bone metabolism. Biochemical markers of bone formation and resorption. Metabolic bone. Procedures for measuring biochemical markers of mineral metabolism.

ACID-BASE BALANCE. Homeostasis acid-base: blood pH buffers. Origin of pH variations. Compensatory mechanisms. Determination of pH, $p\text{CO}_2$ and $p\text{O}_2$. Alterations of the acid-base balance. Respiratory and metabolic acidosis and alkalosis.

Theme III. Biochemical assessment of the function of organs and systems

LIVER FUNCTION. Hepatobiliary system. Liver functions. Catabolism of hemoglobin. Hepatobiliary disease. Laboratory tests for evaluation. Investigation of jaundice: bilirubin determination.

CARDIAC AND MUSCULAR FUNCTIONS. Myocardial infarction and angina. Mechanisms. Meaning diagnostic procedures and measuring creatine kinase and its isoenzymes, myoglobin and troponin. Natriuretic peptides in the diagnosis of heart failure. Myopathies: progressive muscular dystrophy, rhabdomyolysis and polymyositis.

RENAL FUNCTION. Formation of urine and renal functions. Kidney disease: glomerulonephritis, tubular diseases, kidney failure, diabetic nephropathy, renouretal lithiasis. Laboratory tests for evaluation: urea, creatinine, urate, clearance testing, protein and kidney stones.

THYROID FUNCTION. Thyroid. Synthesis, transport, metabolism and regulation of thyroid hormones. Hypothyroidism and hyperthyroidism.

GONADAL AND GESTATIONAL FUNCTIONS. Ovarian function. Hormonal studies: prolactin, estradiol, progesterone, androgen and gonadotropins. Evaluation of infertility. Testicular function. Alterations: hypogonadism, infertility, disorders of puberty. Diagnosis and monitoring of pregnancy. Evaluation of fetus-placental unit.

BIOCHEMISTRY OF CANCER AND TUMOR MARKERS. Concept and classification. Clinical utility. Dynamic interpretation of the results. Main markers: CEA, AFP, b-hCG, PSA, CA19.9, CA125 and CA15.3. Application in different types of tumors. Concept of hereditary cancer. Oncogenes and tumor suppressor genes.

LABORATORY PRACTICES

Session 1

Measures by molecular absorption spectrometry in an analyzer: (1) Measurement of the substance concentration of cholesterol in serum through an enzymatic-colorimetric reaction to an end-point. (2) Measurement of the substance concentration of urea in serum by a two points enzymatic reaction-spectrometric. (3) Measurement of the catalytic activity of L-lactate dehydrogenase (LD) in serum using a continuous spectrometric method recommended by the SEQC^{ML} (Spanish Society of Medical Laboratory). Measuring the substance concentration of total bilirubin in serum through a chemical reaction by molecular absorption spectrometry: Measurements with blank sample. Implementation of an external quality assessment program using the obtained results.

Session 2

Measurement of the catalytic activity of L-lactate dehydrogenase (LD) in serum using a continuous

spectrometric method recommended by the IFCC (International Federation for Clinical Chemistry and Laboratory Medicine) in a molecular absorption spectrometer: Comparison of two procedures and influence of the measuring instrument.

Session 3

Study of the effect of interfering substances (hemoglobin, bilirubin and lipids) in the measurement of the substance concentration of uric acid in serum using an end-point enzymatic procedure in a molecular absorptionspectrometer.

Session 4

Determination of the linearity and the latency periods of the enzymatic reaction catalyzed by aspartate aminotransferase (AST) using a continuous spectrometric method: Effect of pyruvate in the measurement of the enzyme.

Learning activities and methodology

Title	Hours	ECTS	Learning outcomes
Solving problems and practical cases	21	0.84	CM19, CM20, KM29, SM29
Lectures	36	1.44	KM29, SM29
Seminars	10	0.4	CM19, CM20, SM29
Individual study	51	2.04	CM19, CM20, KM29, SM29
Laboratory practices	9	0.36	CM19, SM29
Tutorials	2	0.08	CM19, CM20, KM29, SM29
Evaluation (exams and oral presentations)	10	0.4	CM19, CM20, KM29, SM29

The teaching methodology consists of lectures, seminars and laboratory practices. The teaching materials for these activities can be found on the Virtual Campus.

Lectures: taught in master classes where students acquire basic knowledge of the subject by attending classes and supplementing them with individual study of the topics explained. The lectures are for the whole group.

Seminars: discuss, develop and present case studies, clinical cases and problems. The knowledge acquired in the lectures and individual study will be applied to solve cases and problems. Students work in small groups supervised by the teacher, solving proposed cases and after exposing them to class. The aim of the seminars is to promote the capacity for analysis and synthesis, the critical reasoning and the ability to solve cases and problems.

Laboratory practices: practical sessions in the laboratory where they apply the knowledge acquired in the lectures and promote learning practical techniques of clinical biochemistry laboratory. It promotes teamwork and active self. Students work in small groups. This activity is mandatory. To attend the sessions the student must have passed the biosafety and safety tests found in the Virtual Campus and to be aware and accept the rules of operation of the laboratories of the Faculty of Biosciences.

In this subject, the use of Artificial Intelligence (AI) technologies is allowed as an integral part of the

development of the work, provided that the final result reflects a significant contribution of the student in the analysis and personal reflection. The student must clearly identify which parts have been generated with this technology, specify the tools used and include a critical reflection on how these have influenced the process and the final result of the activity. The lack of transparency in the use of AI will be considered a lack of academic honesty and may lead to a penalty in the grade of the activity, or greater sanctions in serious cases.

Additionally students can have specific tutorials.

Annotation: within the schedule set by the centre or degree programme, 15 minutes of one class will be reserved for students to evaluate their lecturers and their courses or modules through questionnaires.

Assessment

Continuous assessment activities

Title	Weight	Hours	ECTS	Learning outcomes
Clinical cases	15 %	2	0.08	CM20, KM29, SM29
Problems test	10 %	1	0.04	CM20, SM29
2d Midterm exam	30 %	3	0.12	CM19, CM20, KM29, SM29
1t Midterm exam	30 %	3	0.12	CM19, CM20, KM29, SM29
Laboratory practice	15 %	2	0.08	CM19, CM20, KM29, SM29

Evaluation activities

The assessment of the course is continuous and will be evaluated both individually and in group activities. All assessment activities are obligatory. None of the assessment activities represent more than 50% of the final grade.

(1) 1st Midterm exam (30 % of the final grade), written tests on basic concepts of the course. The type will be multiple choice questions (50 %) and short questions (50 %) of the subjects developed in the lectures. It is necessary that the mark of the test is ≥ 4.7 to overcome it. Not showing up the test or failure to pass the test will not allow the student to sit the second midterm exam and they will have to sit the re-assessment exam for the entire subject directly.

(2) 2^d Midterm exam (30 % of the final grade), written tests on basic concepts of the course. The type will be multiple choice questions (50 %) and short questions (50 %) of the subjects developed in the lectures. It is necessary that the mark of the test is ≥ 4.7 to overcome it.

(3) Test problems (10 % of the final grade), evaluation of the problem classes. There is no possibility of re-assessment.

(4) Clinical case (15 % of the final grade), consists of two parts: presentation of the case (10 %) in group of two or three students or individually, content and oral presentation and writing is evaluated; the clinical case presented must be submitted through the Virtual Campus in the deadline. The cases presented but not submitted are not evaluated. To get the remaining 5 % the student must participate by asking questions about the cases presented in class and at the request of the teacher. There is no possibility of re-assessment.

(5) Laboratory practices (15 % of the final grade), evaluation of the practical results obtained in the laboratory

sessions (5 %) and a final written test (10 %). The practices are mandatory. There is no possibility of re-assessment.

Re-assessment exam (60 % of the final grade), for those students who have not passed the midterm exams or who want to mark up. The examination will be the entire subject matter (no individual midterm re-assessment). The type will be multiple choice questions (50 %) and short questions (50 %) of the subjects developed in the lectures and seminars. It is necessary that the mark is ≥ 4.7 to overcome it. To be eligible for the retake process, the student should have been previously evaluated in a set of activities equaling at least 2/3 of the final score of the course or module.

Single assessment

Students who take the single assessment must do the laboratory practices and their assessment in face-to-face sessions. The presentation of the clinical case in person on the assigned day will also be mandatory attendance (attendance on the other seminar days is not mandatory).

The assessment and the weight on the final grade will be the same as those of the continuous assessment (15 % laboratory practices and 15 % presentation of the clinical case).

The single assessment consists of a single summative test (with multiple-choice questions, short questions and problems) on the content of the entire theory program and the program of problem-based seminars.

The grade obtained in the synthesis test is 70 % of the final grade of the subject, the grade obtained in the laboratory practices 15 % and the presentation of the clinical case the remaining 15 %.

The single assessment exam will coincide with the same date fixed in the calendar for the last continuous assessment exam (2nd midterm) and the same recovery system will be applied as for the continuous assessment.

You must consider:

To pass the course you must get at least ≥ 5.0 in the final mark. The non-execution of any of the assessment activities is a zero in that activity.

Total or partial plagiarism from any other source in any assessed activity will automatically be considered as a fail.

The student will be graded as "No Avaluable" if the weight in of all conducted evaluation activities is less than 50 %.

Repeating students keep marks of the test problems, clinical case and laboratory practices activities evaluated for the next academic course. If students do not pass the course in this period, they must return all evaluation activities.

Any irregularity committed during an assessment activity (academic misconduct, plagiarism, or improper use of AI, unless such use is expressly authorized in the course syllabus) that may lead to a significant alteration of the grade will result in that activity being graded as 0. If the course syllabus stipulates that obtaining a minimum mark in this assessment is an essential requirement to pass the course, or if multiple irregularities occur in the assessment activities of the same course, the final grade for the course will be 0. Furthermore, disciplinary proceedings may be initiated against any student who incurs any of these irregularities.

Bibliography

Books

- BIOQUÍMICA CLÍNICA Y PATOLOGÍA MOLECULAR. X Fuentes Arderiu, MJ Castiñeiras Lacambra, JM Queraltó Compañó. Volumes I and II, 2nd ed. Editorial Reverté: Barcelona, 1998
- CLINICAL BIOCHEMISTRY. A Gaw, R Srivastava. Netherlands: Elsevier health Sciences, 2013 (e-book)
- CLINICAL CHEMISTRY. WJ Marshall, M Lapsley. Edinburgh: Elsevier, 2016 (e-book)
- HENRY'S CLINICAL DIAGNOSIS AND MANAGEMENT BY LABORATORY METHODS. JB Henry. Vols. 1 and 2. Elsevier UK, 2016
- TIETZ FUNDAMENTALS OF CLINICAL CHEMISTRY AND MOLECULAR DIAGNOSTICS. CA Burtis, DE Bruns, eds. 7th ed. Elsevier Saunders: St. Louis, MO, 2015
- TIETZ TEXTBOOK OF CLINICAL CHEMISTRY AND MOLECULAR DIAGNOSTICS. N Rifai, AR Horvath and CT Wittwer, eds. 6th ed. Elsevier: St. Louis, MO, 2018 (reference book)

Web pages related to the clinical laboratory (see the Virtual Campus)

Software

There is no specific software for this course

Course groups and languages

The information provided is provisional until November 30. After this date, you will be able to consult the language of each group through this [link](#). To access the information, you will need to enter the course CODE