

Degree programme	Type	Course
Biochemistry	OB	2

Contact lecturer

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Group languages

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

Prerequisites

There are no official prerequisites. However, in order to ensure the proper follow-up of the subject by the student and the achievement of the learning outcomes proposed, it is recommended that the student would have acquired solid knowledge of the following subjects of the 1st year: Organic Chemistry Of the Biochemical Processes, Foundations of General Chemistry, Cell Biology, Biochemistry I and Basic Instrumental Techniques.

On the other hand, in a scientific discipline such as Chemical and Protein Engineering, where many of the sources of information, or at least the most up-to-date ones, are in English, it is highly recommended that students have some basic knowledge of this language .

Objectives

The subject Chemical and Protein Engineering is part of the subject "Molecular Biology", in which the structural and functional characteristics of amino acids and proteins are studied from a basic and applied point of view. Since proteins constitute the effector molecules of many biochemical and biological processes, knowledge of its structure and function is essential for the understanding of a good number of subjects in the Degree in Biochemistry. The theoretical knowledge acquired in the subject of Protein Chemistry and Engineering is complemented by a practical training in the laboratory in the context of Integrated Laboratory 3.

The training objectives are: that the student, at the end of the subject, would be able to:

- Know the chemical and structural properties of amino acids.
- Describe the methods of sequencing and synthesis of peptides.
- Describe the elements of secondary, tertiary and quaternary structure of the proteins, the determinants of their stability and folding.
- Structurally classify proteins.
- Explain the different methods for the determination of the three-dimensional structure of the proteins.
- Describe the molecular bases of the folding of proteins, their molecular dynamics, their post-translational processing and their traffic to the different cellular compartments.
- Explain the biochemical bases of the evolution of proteins.

- Describe the properties of the human proteome and the methods used for its characterization.
- Know the methods for artificial production, modification and optimization of the properties of the proteins
- Integrate and apply the theoretical knowledge acquired to interpret the results of scientific experiments and to solve experimental problems.
- Use of the appropriate scientific terminology.

Learning outcomes

- KM22 (Describe the biochemical and molecular basis of protein folding, intracellular trafficking, post-translational modification, and turnover.) Describe the biochemical and molecular basis of protein folding, intracellular trafficking, post-translational modification, and turnover.
- SM18 (Apply computing resources to visualize and understand the three-dimensional structure of proteins, search for information in databases, and use molecular tools.) Apply computing resources to visualize and understand the three-dimensional structure of proteins, search for information in databases, and use molecular tools.
- SM19 (Analyse the molecular mechanisms that regulate the function of proteins and nucleic acids, as well as their alterations in cancer.) Analyse the molecular mechanisms that regulate the function of proteins and nucleic acids, as well as their alterations in cancer.
- SM20 (Apply the basic techniques of recombinant DNA technology to molecular biology and protein engineering.) Apply the basic techniques of recombinant DNA technology to molecular biology and protein engineering.

Contents

I. FUNDAMENTAL PROPERTIES OF AMINO ACIDS AND PROTEINS.

Proteins, peptides and their functions for living organisms. Structure and physico-chemical properties of amino acids. Chemical reactivity. Differential contribution of the amino acids to the properties of the proteins.

Evolutionary relationships between amino acids.

II. PEPTIDE BOND AND POLYPEPTIDIC SEQUENCE.

Stereochemistry of the peptide bond. Types of natural peptides. Chemical reactivity of peptides. Structural and functional implications of the polypeptide sequence. Strategies for the determination of the protein sequence. Chemical synthesis of peptides;

Combinatorial libraries

III. CONFORMATIONAL DETERMINANTS.

SECONDARY STRUCTURES

Levels of three-dimensional structure. Types of stabilizing forces for the protein conformation. Cooperativity of weak interactions. Conditioners of protein folding. Main types of secondary structures;

\>Amino acids that participate.

\>IV. STRUCTURAL CLASSIFICATION OF PROTEINS

Supersecondary structures and motifs. Structural domains. Tertiary structure. Proteins α . Proteins α / β . Proteins β . Classification methods.

\>Conformation and function in fibrous proteins: α -keratin, fibroin, collagen.

V.

\>FOLDING AND CONFORMATIONAL DYNAMICS.

Protein folding and unfolding: native state and unfolded state. Methods of analysis of folding. Thermodynamic and mechanical characteristics of the folding process. Models that describes folding. Folding and aggregation; Conformational diseases. Protein folding in vivo: molecular chaperones. Proteostasis.

\>Molecular protein dynamics.

VI.

\>POST-TRANSLATION PROCESSES AND MODIFICATIONS.

Types of post-translation modifications and functional implications. Glycosylation. Transport associated modifications. Limited proteolysis: pre-proteins, zymogens. Systems regulated by limited proteolysis: blood clotting, digestive proenzymes.

\>Degradation and half protein life.

\>VII.ENGINEERING OF PROTEINS: REDESIGN AND SYNTHESIS DE NOVO.

Rational design: targeted mutagenesis as a tool for analyzing and modifying proteins. Examples and applications of protein engineering in the analysis of structure, stability, and functionality. Modification and improvement of the properties of proteins. Targeted evolution: protein engineering by combinatorial methods. Examples of recombinant proteins.

\>Protein design de novo.

\>PROBLEMS

The content of this section, which will be presented in the form of a dossier at the beginning of the semester, consists of a determined amount of problems related to the topics developed in Theory. The own characteristics of the various parts of the Theory's program make the problems to concentrate on certain specific aspects that are: properties of amino acids, protein sequencing, protein stability and three-dimensional structure of proteins.

Learning activities and methodology

Title	Hours	ECTS	Learning outcomes
Group tutorials	4	0.16	KM22, SM18, SM19
Theory classes, seminars	37	1.48	KM22, SM18, SM19, SM20
Problems solving	8	0.32	KM22, SM18, SM19
Seminars preparation	7	0.28	KM22, SM18, SM19
Search for information and information management in the process of self-learning (group)	24	0.96	KM22, SM18, SM19, SM20
Study autonomous work	60	2.4	KM22, SM18, SM19, SM20

The training activities are divided into three sections: theory classes, problem classes and seminars each with their specific methodology. These activities will be complemented by a series of tutoring sessions that will be programmed additionally.

Theory classes

The teacher will explain the content of the program with the support of audiovisual material that will be available to the students in the Virtual Campus of the subject in advance at the beginning of each of the subjects of the course.

These lectures will be the most important part of the theory section. It is recommended that students have the material published on the CV in printed form so that they can follow the classes more comfortably.

The tutor would advise the student about the strategies to follow in their learning.

Problem classes

The group will be divided into two subgroups of about 30 students, whose lists will be made public at the beginning of the year. Students will attend the sessions programmed for their group.

At the beginning of the semester a dossier of problems will be presented through the Virtual Campus. They will be resolved throughout the sessions.

Students will work out problems outside class hours. Non-expositive classroom sessions will be devoted to the resolution of previously worked problems during the previous week, which will be discussed and corrected with

the participation of all students.

Seminars

At the beginning of the semester a proposal of subjects will be presented through the Virtual Campus on which students aggrouped in 2-4 groups will be able to prepare a seminar. The difficulties that arise on this self-study material and other issues / problems would be discussed in tutoring classes. The tutor will advise students on the strategies to follow in their elaboration.

Tutorials

These will be carried out with students divided into the same subgroups of problem classes. Its programming will be announced at the beginning of the semester. The purpose of these sessions is to resolve doubts, revise concepts with a high conceptual difficulty and carry out debates on the subjects for which autonomous learning is programmed. These sessions will not advance the subject of the official agenda, but instead will be sessions of debate and discussion.

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
Problem solving in class and seminars	5%	4	0.16	KM22, SM18, SM19
Seminaris	10%	2	0.08	KM22, SM18, SM20
Partial theory tests (individual assessment)	75%	2	0.08	KM22, SM18, SM19
Problem Testing (Individual Assessment)	10%	2	0.08	KM22, SM18, SM19

Theory

The total weight of the evaluation of the theoretical part will be 75% of the total grade of the subject. The main evaluation of this part of the subject will have the format of a continuous evaluation with two partial tests, with another recuperation exam that allows to examine the content of each one of the two partials not previously surpassed, or both simultaneously, if you do not pass any of the partial ones.

The objective of the continuous evaluation is to encourage the student's ongoing effort throughout the course, allowing him to be aware of the degree of follow-up and understanding of the subject.

Students who have passed the partial exams with a note greater than 3.5 on a total of 10 points can choose to obtain the average theory note of the two partials. Those who have not surpassed the value of 3.5 in either partial will have to take the final examination for the subject of the partial or partials in question. In this case, the qualification of the last partial examination will be used to calculate the final grade of theory.

The specific weight of the set of these two tests, or the recuperation exam, is 75% of the total of the mark of the subject.

Problems

The weight of the evaluation of problems will be 20% of the total.

Mixed group / individual assessment:

- Resolution of the problems worked throughout the course and evaluated by the teacher (7.5%)
- Written proof of problems on the date assigned for the final exam of the subject (12.5%). Students who have passed the partial exams with a note greater than 3.5 out of 10 points can choose to obtain the average of the two partial exam notes. Those who have not surpassed the value of 3.5 of either partial will have to take the final examination for the subject of the partial or partials in question; in this case, the qualification of the last examination partial is the one that will be taken to calculate the final grade of problems.

Seminars

The overall assessment of the subject's participation in seminars will count for up to 5% of the total. Thematic seminars may be proposed as a complement to the subject and will always focus on topics not directly addressed in the classroom or the program. If, for logistical reasons, these seminars cannot be held, they will be replaced by research seminars delivered by the course instructor, collaborators, or invited researchers. The content of these seminars will then be assessed as part of the theory examination, which will consequently account for 80% of the final course grade.

Global evaluation:

- The subject will be passed when the sum of the different parts weighted by their specific weight in the subject exceeds 5.0 on 10 points.

To be eligible for the retake process, the student should have been previously evaluated in a set of activities equaling at least two thirds of the final score of the course or module. Thus, the student will be graded as "No Avaluable" if the weighting of all conducted evaluation activities is less than 67% of the final score

Single assessment:

The single assessment consists of a single synthesis test that includes the contents of the entire theory syllabus with a weight of 75% and another of questions corresponding to the SEM (5%), PAUL (20%) with a total weight of 25%. The mark obtained in this synthesis test is 100% of the subject's final mark.

The single assessment test will be held on the same date set in the calendar for the last continuous assessment test, and the same recovery system will be applied as for the continuous assessment.

General considerations:

Any irregularity committed during an assessment activity-including academic fraud, plagiarism, or the improper use of Artificial Intelligence (unless explicitly authorized in the course syllabus)-that may lead to a significant alteration of the assessment outcome will result in a grade of 0 for that assessment.

If the course syllabus specifies that passing this assessment is a mandatory requirement for passing the course, or if multiple irregularities are detected in different assessment activities within the same course, the

student's final course grade will be 0.

In addition, the University may initiate disciplinary proceedings against any student involved in such misconduct.

Use of AI

The use of Artificial Intelligence (AI) technologies in this course is permitted exclusively for support tasks, such as literature and information searches, text editing, and translation, as well as for any specific activities or situations explicitly authorized by the instructor.

Students must clearly identify which parts of their work have been generated or assisted using AI technologies, specify the tools employed, and include a critical reflection on how these technologies influenced both the preparation process and the final outcome of the assignment.

Failure to disclose the use of AI in assessed coursework will be considered a breach of academic integrity and may result in a partial or complete reduction of the assignment grade, or more severe disciplinary sanctions in cases of serious misconduct.

Bibliography

Basic

(from oldest to newest)

- Brandén C. & Tooze J., Introduction to Protein Structure (1999) Garland Science
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Complementary

- Buckel, P. (ed), Recombinant Protein Drugs (2001), Birkhäuser Verlag
- Creighton T.E., Proteins. Structures and Molecular Properties. (1993) (2nd ed.) Freeman W.H. & Co.
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- Patthy, L. Protein Evolution (2007) (2nd ed.) Wiley
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- Park, S.J., Cochran, J.R. Protein Engineering and design (2009) CRC Press
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- Tompa, P. & Fersht, A. Structure and function of intrinsically disordered proteins (2009) CRC Press
- Twyman, R., Principles of Proteomics (2004) Taylor & Francis
- Veenstra, T.D. & Yates, J.R. Proteomics for Biological Discovery (2006) Wiley

Internet links:

You can find them at : https://catalegclassic.uab.cat/search*cat/r?SEARCH=100857

Software

FoldIt

<https://fold.it>

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

Type of teaching	Group	Language	Semester	Shift
(TE) Theory	32	Catalan	first semester	afternoon
(PAUL) Classroom practices	321	Catalan	first semester	morning-mixed
(PAUL) Classroom practices	322	Catalan	first semester	morning-mixed