



Core Bioinformatics

Code: 42397 ECTS Credits: 12

Degree	Туре	Year	Semester
4313473 Bioinformatics	ОВ	0	1

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Teachers

Antoni Barbadilla Prados

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Raquel Egea Sánchez

Xavier Daura Ribera

External teachers

Cedric Notredame

Emanuel Raineri

Josep Abril

Sebastián Ramos

Prerequisites

Level B2 of English or equivalent is recommended.

Objectives and Contextualisation

This module focuses on the development of diverse bioinformatic tools and resources commonly used in Omics research. Our intention is that it covers several aspects of bioinformatics in a series of brief topics, in the form of "tastings". Therefore, it is not an accumulative module, but a transversal one, which should provide

Use of languages

Principal working language: english (eng)

with a wide range of ideas and approaches that bioinformatics offers, through the hands of experts. The main objective is to provide students with the necessary foundation to apply bioinformatics to different areas of scientific research. Over time, each student will be able to gain all the depth they propose on any of these topics, the one which finally represents their research framework.

Skills

- Analyse and interpret data deriving from omic technology using biocomputing methods.
- Design and apply scientific methodology in resolving problems.
- Possess and understand knowledge that provides a basis or opportunity for originality in the development and/or application of ideas, often in a research context.
- Propose biocomputing solutions for problems deriving from omic research.
- Propose innovative and creative solutions in the field of study
- Student should possess the learning skills that enable them to continue studying in a way that is largely student led or independent.
- Understand the molecular bases and most common standard experimental techniques in omic research (genomics, transcriptomics, proteomics, metabolomics, interactomics, etc.)
- Use and manage bibliographical information and computer resources in the area of study
- Use operating systems, programs and tools in common use in biocomputing and be able to manage high performance computing platforms, programming languages and biocomputing analysis.

Learning outcomes

- 1. Create and promote algorithms, calculation and statistical techniques and theories to resolve formal and practical problems deriving from the handling and analysis of biological data.
- 2. Design and apply scientific methodology in resolving problems.
- 3. Identify and apply algorithms in which the programs are based bioinformatic analysis.
- 4. Identify and classify the principle types of biomolecular data obtained from omic technology.
- 5. Possess and understand knowledge that provides a basis or opportunity for originality in the development and/or application of ideas, often in a research context.
- 6. Propose innovative and creative solutions in the field of study
- 7. Search for specific bioinformatics tools and bioinformatics resources in the network.
- 8. Student should possess the learning skills that enable them to continue studying in a way that is largely student led or independent.
- 9. Synthesise and interpret in a logical and reasoned manner the information from the molecular data bases and analyse it using biocomputing tools.
- 10. Understand the theoretical, statistical and biological bases, in which the programs are based bioinformatic analysis: sequence alignment, similarity search and multiple alignment, structure prediction, genome annotation, phylogenetic and evolutionary analysis.
- 11. Use and manage bibliographical information and computer resources in the area of study
- 12. Use the main molecular databases, the main standard formats of molecular data and integrate data from different data sources

Content

Statistical Inference

Professor Antonio Barbadilla

- Statistics: bridge between data and models
- Data Types
- Population and sample
- Experimental design
- Data Quality
- Exploration of Data
- Sample distribution and law of large numbers
- Statistical inference

- Central Limit Theorem
- Point estimation
- Estimation of confidence interval
- Hypothesis
- Elements of a test: H0, H1, statistical test, p value, significance level, type I and II errors, power
- Z test, t test, chi-square test, correlation test, regression, analysis of variance
- Interpretation of statistical significance
- Parametric versus nonparametric tests
- Selecting the appropriate statistical test (decision tree)
- Multivariate Testing
- Resampling

Statistics and Stochastic Processes for Sequence Analysis

Professor Pere Puig

a. Probability basics

Sets and events. Properties. Conditional probability. Independence. Alphabet and sequences. Probabilistic models.

b. The multinomial model

Simulating a multinomial sequence. Estimating probabilities.

c. The seginr package

d. Markov chain models

Concept and examples. Classification of states. R code. Simulating a Markov chain sequence. Estimating the probabilities of transition. The probability of a sequence. Using Markov chain for discrimination.

e. Higher order Markov chain models

Concept and examples. Estimating the probabilities of transition. Comparison of higher order Markov chains.

f. Hidden Markov chain models

Concept and examples. Parameter estimation. Hidden states estimation.

g. An introduction to Generalized Linear Models

GLM basics. The Logistic model. The Poisson model.

Bayesian Inference

Professor Emmanuele Raineri

- 1. Curve fitting.
- Estimation of parameters of probability distributions: binomial, poisson and gaussian.
- Example: fitting a noisy dataset.
- Cross validation, overfitting and regularization.
- 2. Dimensional reduction.
- Principal component analysis, multidimensional scaling.
- Example: distinguishing cell types using methylation profiles.
- 3. Lasso regression.
- Variable selection in linear models.
- Penalized regression: Lasso and Elastic Net.
- Example: lasso regression in R.

Bioinformatics Formats and Databases

Professor Daniel Yero

a. Sequence formats

Nomenclature. Text editors. FASTA format and its variants. Raw/Plain format. Genbank sequence format. EMBL sequence format. GCG, NBRF/PIR, MSA, PHYLIP, NEXUS. Format conversion.

b. Databases

Concept. Boolean searches. Wildcards and regular expressions. Identifiers and accession numbers. Classification. NAR databases compilation. GenBank and other NCBI databases. EMBL. DDBJ. Integrated Meta-Databases. Main nucleotide, protein, structure, taxonomy, etc. databases.

Workflows with Galaxy

Professor Raquel Egea

a. Introduction to workflow managers

Concept, origin and design of workflow managers. Workflow patterns. Existing workflow managers. APIs and Web Servers.

b. Galaxy: basics, interface and practical uses.

Software Engineering

Professor Miquel Angel Senar

- a. Parallelization strategies and HPC
- b. Cloud computing with Amazon Web Services
- c. Version control system with Git and GitHub

Sequence Alignment

Professor Cedric Notredame

a. Evolution and comparison Models

Molecular clock. Protein structure and evolution. Substitution Matrices.

b. Dynamic Programming based sequence comparisons

Needlman and Wunsch algorithm. Smith and Waterman algorithm. Afine gap penalties computation. Linear space computation of pairwise algorithms.

c. Blast and Database searches

The Blast algorithm. E-values and estimates of statistical significance. Database search strategies. PSI-Blast and other evolutionary approaches.

d. Multiple Sequence Alignments: algorithms and strategies

Main applications of multiple sequence alignments. Most common algorithms. Multiple sequence alignment strategies.

Gene and Control Region Finding

Professor Josep Abril

a. Gene prediction

Annotation: concept, databases, problems. Gene finding: search by signal, search by content, approaches (ab-initio, homology search, comparative genomics, NGS), evaluation of software accuracy.

b. Finding regulatory motifs

DNA motifs: exact matching, regular expressions, position weight matrices, search trees, profiles, randomized algorithms, logos and pictograms, software for motif finding. Regulatory domains. Histones. CRMs architecture & networking. Regulatory networks. Meta-alignment. Conservation, phylogenetic footprinting and phylogenetic shadowing. NGS.

Population Genomics

Professor Alfredo Ruiz

a. Population genomics under neutrality in a finite population

Introduction. Genetic drift. Effective population size. Probability of fixation of neutral mutants.

b. Population genomics under selection

Natural selection. Probability of fixation of selected mutants. Fitness distribution of new mutants. Rate of evolution.

c. Adaptive evolution and population size

Phylogeny and Molecular Evolution

Professor Sebastián Ramos

a. Models of sequence evolution

DNA sequence. Jukes and Cantor model. More realistic models. Model selection.

b. Phylogeny

Concept. Species trees versus gene trees. Tree-reconstructionmethods: distance methods, maximum parsimony, maximum likelihood, Bayesian inference. Support. Phylogenomics. Building trees with R.

Introduction to Genomes

Professor Alfredo Ruiz

a. Introduction to genomes

Sequenced genomes. Organization and size of eukaryotic genomes. Building a genome: NGS methods for genomics and transcriptomics.

b. The human genome: where are we now?

Current assembly of the human genome. The ENCODE project: functional elements in the human genome. Repetitive content of the human genome.

Structural Bioinformatics

Professors Xavier Daura, Óscar Conchillo, Leonardo Pardo and Jean-Didier Maréchal

a. Structural biology and interactions

Biomolecules. Xenobiotics. Proteins, aminoacids and peptide bonds. Four levels of protein structure. Protein folding and stability. DNA structure. Experimental methods for structure determination.

b. Structure databases

PDB. PyMOL.

c. Molecular modeling

Models. Potential energy. Quantum and molecular mechanics techniques. Energy minimization. Molecular Dynamics. Homology modeling.

d. Docking

Concept and applications. Exploring the conformational space. Defining molecular interactions. Scoring functions. Workflows and software. Online resources.

Systems Biology

Professor Isaac Salazar

a. Classical and Genomic age Systems Biology

The systems biology paradigm in light of technological developments over the last 100 years. Data integration bottlenecks.

b. Mathematical modeling of molecular circuits.

Conceptual models. From conceptual models to mathematical models. Mathematical formalisms. Data driven models.

c. Design and organization principles in molecular circuits.

Concept of design principle. Mathematically controlled comparisons. Feasibility analysis. Design Spaces. Synthetic Biology.

Methodology

The methodology will combine master classes, solving practical problems and real cases, working in the computing lab, performing individual and team work, reading articles related to the thematic blocks, and independent self-study. The virtual platform will be used.

Activities

Title	Hours	ECTS	Learning outcomes
Type: Directed			
Solving problems in class and work in the biocomputing lab	39	1.56	
Theoretical classes	39	1.56	
Type: Supervised			
Performing individual and team works	40	1.6	
Type: Autonomous			
Regular study	178	7.12	

Evaluation

The evaluation system is organized in three main activities. There will be, in addition, a retake exam. The details of the activities are:

Main evaluation activities

- Student's portfolio (55%): work done and presented by the student all along the course. None of the individual assessment activities will account for more than 50% of the final mark.
- Individual theoretical and practical test (35%): a final exam will take place at the end of this module.

 It will consist of one or two multiple-choice or short questions by each professor teaching in this module.
- Soft skills (10%): assistence, arrival on time and active participation in class.

Retake exam

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 module. The teacher will inform the procedure and deadlines for the retake process. Please note that soft skills cannot be recuperated.

Not valuable

The student will be graded as "Not Valuable" if the weight of the evaluation is less than 67% of the final score.

Evaluation activities

Title Weighting Hours	ECTS Learning outcomes
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Individual theoretical and practical test	35%	4	0.16	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12
Soft skills	10%	0	0	2, 5, 6, 8
Student's portfolio	55%	0	0	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12

Bibliography

Updated bibliography will be recommended in each session of this module by the professor, and links will be made available on the Student's Area of the MSc Bioinformatics official website (http://mscbioinformatics.uab.cat/base/base3.asp?sitio=bioinformaticsintranet&anar=module_2 &item=&subitem=).