		Teachin	g Guide			
Identifying Data				2016/17		
Subject (*)	Bioinformática e Modelado de Biomoléculas Code			610441020		
Study programme	Mestrado Universitario en Bioloxía Molecular , Celular e Xenética					
		Descr	iptors			
Cycle	Period	Ye	ear	Туре	Credits	
Official Master's Degree	2nd four-month period	Fir	rst	Optativa	3	
Language	Spanish					
Teaching method	Face-to-face					
Prerequisites						
Department	Bioloxía Celular e MolecularComp	utaciónTecnol	oxías da Informació	ón e as Comunicación	s	
Coordinador	Dorado de la Calle, Julian		E-mail	julian.dorado@u	dc.es	
Lecturers	Becerra Fernandez, Manuel		E-mail	manuel.becerra	@udc.es	
	Dorado de la Calle, Julian			julian.dorado@udc.es		
Web						
General description	Knowledge management in biology is the field of bioinformatics, and includes both the formalization of the information					
	obtained and its organization in appropriate databases, the extraction of relationships between the scattered information,					
	the modeling of biological processes and the generation of hypotheses to support new experimental approaches. From a					
	technical standpoint, bioinformatics using computational methods (the proper method development in this area is often					
	called computational biology) and receives contributions from mathematics, physics and computer engineering. However,					
	from the point of view of the objectives, bioinformatics is a branch of biology, as they can be biochemistry or microbiology.					
	This interdisciplinary nature of bioinformatics lies both its strength and its weakness: first, the application of ideas brought					
	from other fields consistently produ	uces spectacu	lar advances; but o	n the other hand, it is	difficult to develop appropriate	
	training programs.					
	To realize the importance of bioinformatics in modern biology, it may enough to say that the method most cited publications					
	in this area is Blast, a computational method that searches and identifies sequences of proteins and nucleic acids in					
	databases: ie more technical operations is performed by computational biologists, and no experimental. In fact, the					
	interpretation of any experiment in biology requires complex, almost inevitably, bioinformatic analysis, which is especially					
	obvious in massive experiments.					

	Study programme competences / results
Code	Study programme competences / results
А3	Skills of understanding the functioning of cells through the structural organization, biochemistry, gene expression and genetic variability.
A9	Skills of understanding the structure and dynamics of proteins to individual and proteomic level, as well as the techniques that are
	necessary to analyze them and to study their interactions with other biomolecules.
A11	Skills of understanding the structure, dynamics and evolution of genomes and to apply tools necessary to his study.
B1	Analysis skills to understand biological problems in connection with the Molecular and Cellular Biology and Genetics.
B2	Skills of decision making for the problem solving: that are able to apply theoretical knowledges and practical acquired in the formulation of
	biological problems and the looking for solutions.
В3	Skills of management of the information: that are able to gather and to understand relevant information and results, obtaining conclusions
	and to prepare reasoned reports on scientific and biotechnological questions
B9	Skills of preparation, show and defense of a work.
C3	Skills of Using basic tools of the information technologies and communications (ICT) necessary to the exercise of his profession and for
	the apprenticeship over his life.
C6	Considering critically the knowledge, technologies and the available information to solve problems with which should face.
C8	Considering the importance that the investigation has, the innovation and the technological development in the socioeconomic advance
	and cultural of the society.

Learning outcomes				
Learning outcomes		Study programme		
	con	npetenc	es/	
	results		i	
Know access to Channels Bioinformatics Web Resources	AR3	BR3	CC3	
		BR9		
Understand and manage properly the area of Bioinformatics	AR3	BR3	ССЗ	
		BR9	CC6	
Being able to function independently to find information about the different programs and their changeable parameters and	AR3	BR2	ССЗ	
understand the impact on the results of the analysis		BR3		
		BR9		
To have bioinformatics knowledge of how to make a prediction of the onedimensional characteristics of a protein	AR3	BR1	CC3	
	AR9	BR2	CC6	
	AR11	BR3	CC8	
To be able to perform a simple prediction of the three dimensional structure of a protein based on available data and programs	AR3	BR1	CC3	
on the Web		BR2	CC6	
		BR3	CC8	
Learn the basic methods of molecular simulation and how they are used for the study of proteins	AR3	BR1	ССЗ	
		BR2	CC6	
		BR3	CC8	

	Contents
Topic	Sub-topic
Bioinformatics	Web Resources and Databases in molecular biology. Analysis and comparison of
	sequences.
	Sequence alignment. Location of motives. Search of genes. annotation of
	genes. Browsers genome project. Examples of applications. Data analysis.
Modeling of Biomolecules	Prediction of the characteristics of the protein structure. Obtaining three-dimensional
	models.
	Homology modeling. Modeling by threading or by remote homology design.
	Ab initio methods. Evaluation of the prediction methods.

	Plannir	ng		
Methodologies / tests	Competencies /	Teaching hours	Student?s personal	Total hours
	Results	(in-person & virtual)	work hours	
Guest lecture / keynote speech	A3 A9 A11	10	20	30
Seminar	B3 B9 C6 C8	2	7	9
Laboratory practice	B2 B1 C3	9	22.5	31.5
Personalized attention		4.5	0	4.5
(*)The information in the planning table is for	guidance only and does no	t take into account the I	neterogeneity of the stud	dents.

Methodologies		
Methodologies	Description	
Guest lecture /	Oral presentation complemented by the use of audiovisual media for the purpose of transmitting knowledge and facilitate	
keynote speech	learning.	
Seminar	Working technique that aims to make powerpoint and word documents on a topic proposed by the teacher.	
Laboratory practice	Methodology that allows students to learn effectively through practical activities (demonstrations, simulations, etc.) the theory	
	of a field of knowledge through the use of information technology and communications.	

Personalized attention

Methodologies	Description
Seminar	The personal attention that is described in relation to these methodologies are conceived as moments of classroom student
Laboratory practice	work with teacher, this involve mandatory participation for the student.
	The manner and time in which it was held is indicated in relation to each activity along the course according to the work plan of
	the course

		Assessment	
Methodologies Competencie		Description	
	Results		
Guest lecture /	A3 A9 A11	A test will be realized to assess the knowledge acquired in the course of lectures.	30
keynote speech			
		With this methodology the A5, B2 skillls will be assessed	
Seminar	B3 B9 C6 C8	The seminar will be evaluated by taking into account the ability to extract the most	25
		relevant information obtained for the student, the capacity for teamwork and the ability	
		to expose in public.	
		Whit this methodology B1, B3 and B9 competencies will be evaluated	
Laboratory practice	B2 B1 C3	Regular attendance and active participation in the lab, as well as the bulletin	45
		responses made by students will be assessed. They also perform a test to assess the	
		knowledge acquired.	
		With this methodology the A5 and B2 competencies will be assessed	

Assessment comments

Students presented in the first opportunity of June will be eligible to get honours.

Students with a part-time assistance or exemption may agree with teachers specific methods for evaluation early in the course .

Also students engaged as "SEMIPRESENCIALES" should contact the teachers in the first weeks.

Sources of information

Basic

BIOINFORMÁTICA? Attwood, T.K. & D.J. Parry-Smith. 1999. Introduction to Bioinformatics. Addison Wesley Longman Limited, Edimburgo. ? Baxevanis, A.D. & B.F. Francis Oullette (Eds.). 2002. Bioinformatics. A practical guide to the analysis of genes and proteins. 2nd Ed.Wiley-Interscience.? Bishop, M. 1999. Bioinformatics. Taylor & Francis, UK.? Claverie, J.M. and C. Notredame. 2003. Bioinformatics for dummies. Wiley Publishing, Inc.? Gibas, C. y P. Jambeck. 2001. Developing Bioinformatics Computer Skills. O'Reilly? Higgins, D. y W. Taylor. 2000. Bioinformatics: Sequence, structure and databanks. Oxford University Press.? Higgs, P. & T.K. Attwood 2005. Bioinformatics and molecular evolution. Blackwell Publishing.? Kanehisa, M. 2000. Post-genome informatics. Oxford University Press? Li, W-H. 1999. Molecular evolution. Sinauer Associates Inc., Massachusetts, 2nd. Ed.? Mount, David W. 2001. Bioinformatics. Sequence and Genome Analysis. Cold Spring Harbor Laboratory Press.? Nei, M. y S. Kumar. 2000. Molecular Evolution and Phylogenetics. Oxford University Press.? Pevsner, J. 2003. Bioinformatics and Functional Genomics. John Wiley & Sons, Inc.? Rashidi, H.H. and L.K. Buehler. 2000. Bioinformatics Basics. Applications in Biological Science and Medicine. CRC Press, Boca Raton.? Salzberg, S., D. Searls, and S. Kasif (Eds). 1998. Computational Methods in Molecular Biology. Elsevier Science.? Swindell, S.R., R.R. Miller y G.S.A. Myers. 1997. Internet for the Molecular Biologist. Horizon Scientific Press, Norfolk, UK.? Tisdall, J. 2001. Beginning Perl for Bioinformatics. O'ReillyMODELADO DE BIOMOLÉCULAS? Bnaszak, L. J. 2000. Foundations of structural biology. Academic Press. ? Bourne, P. E., Weissig, H. 2003. Structural Bioinformatics. John Wiley & Sons.? Branden, C. & Tooze, J. 1998. INTRODUCTION TO PROTEIN STRUCTURE. 2nd editionGarland Publishing, Inc, New York . ? Creighton, T. E. 1993. PROTEINS: STRUCTURES AND MOLECULAR PROPERTIES, 2nd edition. W.H.Freeman & Company, New York .? Gómez-Moreno, C. & Sancho, J. (Coords). 2003. ESTRUCTURA DE PROTEÍNAS. Ariel Ciencia, Barcelona . ? Lesk, A.M. 2000. INTRODUCTION TO PROTEIN ARCHITECTURE. THE STRUCTURAL BIOLOGY OFPROTEINS. Oxford University Press, Oxford . ? Tramontano, A. 2006. Protein Structure Prediction. Wiley-Vch.

Complementary

Programas de visualización molecular: Rasmol: http://www.umass.edu/microbio/rasmol Swiss-PdbViewer: http://www.expasy.ch/spdbv/ MOLMOL http://www.mol.biol.ethz.ch/wuthrich/software/molmol Cn3D http://www.ncbi.nlm.nih.gov/Structure/CN3D/cn3d.shtml Chime http://www.umass.edu/microbio/chime Servidores de predicción e modelización: SWISS-MODEL http://expasy.ch/swissmod/ The PredictProtein Server http://www.embl-heidelberg.de/predictprotein/predictprotein.html Center for Molecular Modeling: http://cmm.info.nih.gov/modeling/ GRAMM: http://reco3.musc.edu/gramm/ PQS (Probable Quat. Structure): http://msd.ebi.ac.uk/services/quaternary/quaternary.html

	Recommendations
	Subjects that it is recommended to have taken before
Técnicas Moleculares/610441002	2
	Subjects that are recommended to be taken simultaneously
Dinámica e Estructura de Proteín	as/610441011
Proteómica/610441013	
Xenómica/610441014	
	Subjects that continue the syllabus
Traballo de Máster/610441022	
	Other comments

(*)The teaching guide is the document in which the URV publishes the information about all its courses. It is a public document and cannot be modified. Only in exceptional cases can it be revised by the competent agent or duly revised so that it is in line with current legislation.