

		Teaching Guide				
	Identifying	J Data		2020/21		
Subject (*)	Data structures and algorithmics fo	or biological sequences	Code	614522013		
Study programme	Mestrado Universitario en Bioinforr	Mestrado Universitario en Bioinformática para Ciencias da Saúde				
		Descriptors				
Cycle	Period	Year	Туре	Credits		
Official Master's Degre	e 2nd four-month period	First	Obligatory	6		
Language	SpanishEnglish					
Teaching method	Hybrid					
Prerequisites						
Department	Ciencias da Computación e Tecnol	loxías da InformaciónComputac	ión			
Coordinador	Ladra González, Susana	E-mail	susana.ladra@uc	dc.es		
Lecturers	Ladra González, Susana	E-mail	susana.ladra@ud	.es		
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Web		l	1			
General description	This course introduces the basics of	of the algorithms and data struct	tures that are commonly	used in the field of		
	computational biology.					



Contingency plan	1. Modifications to the contents
	- No changes will be made
	2. Methodologies
	*Teaching methodologies that are maintained
	- Guest lecture / keynote speech
	- ICT practicals
	- Supervised projects
	- Personalized attention
	*Teaching methodologies that are modified
	- Mixed test: in case it cannot be performed in person, it will be replaced by "Problem solving")
	3. Mechanisms for personalized attention to students
	- Email: Daily. Used to make queries, request virtual meetings to resolve doubts and monitor the work being supervised.
	- Moodle: Daily. According to the needs of the students. They will have "thematic forums" associated to the modules of the
	subject, to formulate the necessary consultations.
	- Teams: Telematic session in the time slot assigned to the subject in the faculty's classroom calendar for the advancement
	of the expository and practical subject. Individual or small group telematic sessions to resolve doubts about the subject of
	the presentation, practice or supervised work
	4. Modifications in the evaluation
	The 30% of the mixed test is replaced by "Problem solving", which will also have a 30% weight in the evaluation. It will
	consist of the resolution of four exercises in which the knowledge and the
	competencies acquired during the master classes. The students will have to hand in the solution of the proposed exercises,
	having to defend it orally.
	*Evaluation observations:
	The minimum requirements for the different parts of the evaluation will not be applied. The final mark will be the sum of
	those obtained in each part.
	FIRST OPPORTUNITY:
	Any student who does not send a proposal for the proposed problem-solving tasks will be marked as "Non presentado"
	(Absent).
	SECOND OPPORTUNITY:
	Only those students who do not pass the subject in the first opportunity will be able to present themselves to the second
	opportunity. The recovery of each part will be done in the following way:
	- ICT practicals (50%): students will be able to repeat the proposed ICT practicals during the course in the same conditions
	as in the first opportunity (the internships delivered late will obtain a maximum of 80%). Thus, in case of repeating all the
	practices, the maximum mark that can be obtained is 4 points.
	- Supervised project (20% of the final mark): carried out under the same conditions as at the first opportunity, but on an
	individual basis.
	- Problem solving (30%): carried out under the same conditions as the first time.
	- In case of not recovering any of the parts, the grade obtained at the first opportunity in that part will be kept.

- Any student who does not opt for the recovery of any of the parties will have a grade of "Non presentado" (Absent).



5. Modifications to the bibliography or webgraphy

- No changes will be made.



	Study programme competences / results
Code	Study programme competences / results
A1	CE1 - Ability to know the scope of Bioinformatics and its most important aspects
A2	CE2 ? To define, evaluate and select the architecture and the most suitable software for solving a problem in the field of bioinformatics
A3	CE3 ? To analyze, design, develop, implement, verify and document efficient software solutions based on an adequate knowledge of the
	theories, models and techniques in the field of Bioinformatics
A8	CE8 - Understanding the basis of the information of the hereditary material, its transmission, analysis and evolution
A9	CE9 ? To understand the benefits and the problems associated with the sequencing and the use of biological sequences, as well as
	knowing the structures and techniques for their processing
B1	CB6 - Own and understand knowledge that can provide a base or opportunity to be original in the development and/or application of ideas,
	often in a context of research
B2	CB7 - Students should know how to apply the acquired knowledge and ability to problem solving in new environments or little known within
	broad (or multidisciplinary) contexts related to their field of study
B8	CG3 - Be able to work in a team, especially of interdisciplinary nature
C6	CT6 - To assess critically the knowledge, technology and information available to solve the problems they face to.
C7	CT7 ? To maintain and establish strategies for scientific updating as a criterion for professional improvement.

Learning outcomes			
Learning outcomes		Study programme	
	con	npetenc	es/
		results	
To know the data structures and the algorithms used for compactly storing and processing of biological sequences.	AJ1		
	AJ2		
	AJ9		
To analyze and compare the data structures and the complexity of the algorithms used.	AJ2	BJ1	CJ6
	AJ3		CJ7
To understand, analyze, design and implement solutions for different fundamental problems of sequence alignment, read error	AJ1	BJ1	CJ6
correction, contig assembly, gap filling, etc.	AJ2	BJ2	CJ7
	AJ3	BJ8	
	AJ8		
	AJ9		
To explain, analyze, design and implement solutions to the problems related with evolution, such as haplotype assembly, motif	AJ1	BJ1	CJ6
finding, permutation patterns, genomic rearrangement, etc.	AJ2	BJ2	CJ7
	AJ3	BJ8	
	AJ8		
	AJ9		

Contents		
Topic Sub-topic		
Introduction to algorithms complexity analysis	Algorithms analysis	
	Complexity	
Sequence pattern search	Exact string matching methods	
	Approximate string matching methods	
	Suffix trees and suffix arrays	
Introduction to sequence compression and indexing	Compression techniques	
	Indexes and self-indexes	



Applications to biological sequences	Sequence comparison
	Motif finding
	Genomic rearrangements
	Sequence alignment
	Sequence assembly
	Phylogenetic analysis

	Plannin	g		
Methodologies / tests	Competencies /	Teaching hours	Student?s personal	Total hours
	Results	(in-person & virtual)	work hours	
ICT practicals	A2 A3 B1 B2 B8 C6	14	60	74
	C7			
Supervised projects	A1 A2 A3 A8 A9 B1	3	30	33
	B2 B8 C6 C7			
Mixed objective/subjective test	A1 A2 A3 A8 A9 B2	0	5	5
Guest lecture / keynote speech	A1 A2 A3 A8 A9	28	10	38
Personalized attention		0	0	0

(*)The information in the planning table is for guidance only and does not take into account the heterogeneity of the students.

Methodologies	
Methodologies	Description
ICT practicals	Students will complete practical exercises to develop all the knowledge acquired during lectures.
Supervised projects	Students will develop a work, individually or in small group, under the supervision of the teachers.
Mixed	It consists of a written test to show that the student has acquired the knowledge and skills during lectures and practice
objective/subjective	sessions.
test	
Guest lecture /	Lectures where the course contents are exposed.
keynote speech	

	Personalized attention		
Methodologies	Description		
Supervised projects	There may exist differences among the students regarding their background on algorithms and data structures. Thus, teachers		
ICT practicals	will provide personalized attention for practice sessions and for the supervised project, both individual or in small groups.		

Assessment			
Methodologies	ologies Competencies / Description		Qualification
	Results		
Mixed	A1 A2 A3 A8 A9 B2	It will consist of a written test where the students must prove the knowledge and	30
objective/subjective		competences acquired during lectures and practice sessions.	
test			
		To pass the course globally it is necessary to obtain in the mixed test a minimum	
		grade of 1.5 (over 3). If that minimum grade is not achieved, the maximum grade	
		cannot exceed 4.9 (and therefore the course is failed)	
Supervised projects	A1 A2 A3 A8 A9 B1	Students must complete a project, individually or in small groups, related with a	20
	B2 B8 C6 C7	scientific article. It must be presented in front of the teaching staff.	



ICT practicals

A2 A3 B1 B2 B8 C6 C7 The work done by the students during practice sessions will be assessed. Students must submit bulletins with their solutions to proposed problems and defend them in front of the teaching staff.

Assessment	comments
ASSESSINCIL	COMMENCE

FIRST OPPORTUNITY:

Students that do not take the written exam will obtain a grade of "Non presentado" (Absent).

SECOND OPPORTUNITY:

Only those students that have not passed the course in the first opportunity can be evaluated in the second opportunity. Students can recover any of the parts as follows:

ICT practicals (50%): the students can repeat the ICT practicals under the same circumpstances than in the first opportunity (those submitted out of time can obtain a maximum of 80% of the grade). Thus, in case of repeating all the assignments, the maximum grade will be 4 points. Supervised project (20%): in the same conditions as in the first opportunity. Written test (30%): in the same conditions as in the first opportunity. Written test (30%): in the same conditions as in the first opportunity. In case of not retaking one of the parts, the grade obtained in the first opportunity for that part will be kept. To pass the course globally it is necessary to obtain in the mixed test a minimum grade of 1.5 (over 3). Students that do not retake any part will obtain a grade of "Non presentado" (Absent). ADVANCED OPPORTUNITY:

The assessment for the advanced opportunity is equivalent to that of the first opportunity (50% ICT practicals, 20% supervised project, which must be done individually, and 30% written text). ACADEMIC DISPENSATION:

Students officially enrolled part-time who have been granted an official dispensation from attending classes, as stipulated in the regulations of this University, must contact with the responsible of the course within the first two weeks to establish the conditions for submitting and defending the practical exercises and the supervised project.

	Sources of information
Basic	- Dan Gusfield (1997). Algorithms on Strings, Trees and Sequences. Cambridge University Press
	- Neil C. Jones, Pavel A. Pevzner (2004). An Introduction to Bioinformatics Algorithms. MIT Press
	- Veli Mäkinen, Djamal Belazzougui, Fabio Cunial, Alexandru I. Tomescu (2015). Genome-Scale Algorithm Design.
	Cambridge University Press
Complementary	- Enno Ohlebusch (2013). Bioinformatics Algorithms: Sequence Analysis, Genome Rearrangements, and Phylogenetic
	Reconstruction. Oldenbusch Verlag
	- G. Navarro y M Raffinot (2002). Flexible Pattern Matching in Strings. Cambridge University Press
	- A. Moffat y A. Turpin (2002). Compression and Coding Algorithms. Kluwer Academic Publishers
	- T. C. Bell, J. G. Clearly y I. H. Witten (1990). Text Compression. Prentice Hall

Recommendations
Subjects that it is recommended to have taken before
Introduction to molecular biology/614522004
Genetics and molecular evolution/614522005
Genomics/614522006
Fundamentals of bioinformatics/614522008
Introduction to programming/614522001
Subjects that are recommended to be taken simultaneously
Subjects that continue the syllabus
Advanced processing of biological sequences/614522020
New trends and applications in bioinformatics and biomedical engineering/614522021
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.