POZNAN UNIVERSITY OF TECHNOLOGY



EUROPEAN CREDIT TRANSFER AND ACCUMULATION SYSTEM (ECTS)

COURSE DESCRIPTION CARD - SYLLABUS

Course name Phylogenetic Analysis [S2Bioinf2>AFIL]

Course			
Field of study Bioinformatics		Year/Semester 1/2	
Area of study (specialization) –		Profile of study general academic	:
Level of study second-cycle		Course offered in Polish	
Form of study full-time		Requirements compulsory	
Number of hours			
Lecture 15	Laboratory classe 15		Other (e.g. online) 0
Tutorials 0	Projects/seminars 0	6	
Number of credit points 2,00			
Coordinators mgr Mateusz Twardawa mateusz.twardawa@put.poznan.p	I	Lecturers	

Prerequisites

Students shall know basis of evolutionary processes, molecular biology, algorithms and data structures, machine learning, optimization and statistical data analysis.

Course objective

The main course objective is to equip students with knowledge and skills needed to preform phylogenetic analysis. Students will learn theoretical concepts and practical use of modern methods, algorithms and tools that are crucial in phylogenetic tree reconstruction and analysis.

Course-related learning outcomes

Knowledge:

1. Student possess theoretical knowledge about phylogenetic and is able to use it in order to solve biological problems based on real data.

2. Student understands essential methods and algorithms in phylogenetic analysis, including statistical analysis of phylogenetic trees.

3. Student has knowledge about chosen tools and algorithms that are used in phylogenetic analysis. Student understands phylogenetic techniques and research problems in sequence similarity.

4. Students knows advanced concepts in phylogenetic analysis and is able to use them while planning and designing phylogenetic analysis for different organisms. Student incorporates knowledge about molecular evolution and genomics while solving phylogenetic problems.

5. Student is familiar with databases, tools and modern methods in phylogenetic analysis and knows trends in this discipline.

Skills:

1. Student is proficient in use of selected computer tools in phylogenetic analysis. Student understands phylogenetic techniques and is able to use it properly in sequence similarity analysis. Student can select proper tools, algorithms and methods to solve specific biological problem related to phylogenetics (in scope of this course).

2. Student is proficient in use of statistical methods and tools that perform phylogenetic tree analysis and evaluation.

3. Student is able to analyze phylogenetic problem and pick the best approach to solve it.

4. Student knows how to formulates phylogenetic hypotheses and perform relevant analysis to test them.

Social competences:

1. Student understands the need for lifelong learning and sees practical usage of gained knowledge in phylogenetics (students will learn about phylogenetic analysis of viruses during epidemics and will be presented selected publications about new approaches in phylogenetic analysis).

Methods for verifying learning outcomes and assessment criteria

Learning outcomes presented above are verified as follows:

Formative Assessment:

For Lectures:

The verification of intended learning outcomes is conducted through:

- Responses to questions about material covered in previous lectures.

For Laboratory Sessions:

The verification of intended learning outcomes is conducted through:

- Evaluation of the student's preparation for individual laboratory sessions and their skills in completing planned tasks.

- Continuous assessment during each session (oral responses), emphasizing the progression of skills in applying learned principles and methods.

- The dominant component of formative assessment is the evaluation of reports prepared partially during the sessions and partially afterward.

Summative Assessment:

For Lectures:

Knowledge acquired during the lectures will be verified through a test.

For Laboratory Sessions:

The verification of intended learning outcomes is conducted through the evaluation of reports prepared during individual laboratory sessions. Each task set is scored.

- Students must achieve more than 50% of the points allocated to laboratory sessions to pass.

- All reports must be completed and passed to receive a positive final grade for the laboratory sessions.

- The final grade for the laboratory is based on the total points accumulated across all sessions. Overall Course Grading Scale:

- Below 51%: Fail (2.0)

- 51-60%: Sufficient (3.0)

- 61-70%: Sufficient Plus (3.5)

- 71-80%: Good (4.0)

- 81-90%: Good Plus (4.5)

- 91-100%: Very Good (5.0).

Programme content

Lectures on Phylogenetic analysis can be divided into 7 parts.

First part of the lectures is meant to introduce students to phylogenetics. During this part some crucial problems and needs of phylogenetic analysis will be covered. Moreover, students will be introduced to

elements of biological systematics of prokaryotic and eukaryotic organisms. The importance of reliable systematics, taxonomy and phylogenetics in biological and medical research will be highlighted. Second part of lectures will concentrate on mechanisms guiding evolution on molecular level and genomic diversity. In this part most common difficulties and problems in phylogenetic analysis will be covered such as misuse of biological data and data imperfections or inaccessibility. In scope of this part terms and definitions for crucial concepts will be presented including: homoplasy, homology, use of morphological and other various types of data as phylogenetic analysis support. Methods and criteria for traits and biological sequence proper choice will be given for different problems with phylogenetic analysis close and distant related biological taxa (molecular markers). During this part concept of molecular clock will be presented and problems related to it.

Third part of lectures will cover different types of phylogenetic trees and graph representations of similarity. Moreover, methods for data transformation and processing will be presented as well as natural and artificial taxa and phylogenetic tree rooting.

Fourth part of the lectures will focus on algorithms and tools that are used in phylogenetic tree reconstruction. Students will be introduced to general concept of clustering and data labeling. Chosen and most widely used methods in phylogenetic analysis will be presented, that is distance approaches, parsimony approaches, maximum likelihood approaches and Bayesian approaches. Chosen tools in phylogenetics will be briefly characterized in addition to this part.

The fifth part of lectures will concentrate on statistical analysis of phylogenetic trees and results of analysis. Methods for tree sampling and branch support will be covered as well as terms such as phylogenetic signal, Bremer index, bootstrap, aLTR, posterior probability. In addition some methods for divergent time analysis and consensus tree construction will be presented.

Sixth part of lectures will focus on heuristic approach to phylogenetic analysis and analysis and visualization of results. During this part methods for scoring and validation of phylogenetic trees will be presented.

Last part of lectures will be covering applications of phylogenetic analysis to modern biological research. Chosen methods for phylogenetic analysis of proteins will be presented, but also analysis and monitoring of viruses evolution during epidemics.

Laboratory part of the course will focus on development practical skills in phylogenetic analysis by problem solving. During each laboratory classes students will work with set of tools, algorithms and databases in order to learn how to perform end-to-end phylogenetic analysis. Students will practice different ways to construct, analyze, visualize and score phylogenetic trees. Moreover, student will learn how to monitor evolution of viruses during epidemics.

Course topics

Lecture:

- 1. Introduction to Phylogenetics
- 2. Molecular Evolution Models and Distance Calculation for Biological Sequences
- 3. Molecular Dating and Phylogenetic Trees
- 4. Tree Searching and Probability-based Methods
- 5. Visualization, Analysis, and Evaluation of Obtained Trees
- 6. Applications and Contemporary Trends in Phylogenetics
- Laboratories:
- 1. Tree Construction Based on Multiple Sequence Alignments
- 2. Distance Methods Additive Methods and Distance Matrices
- 3. Distance Methods Ultrametric Trees and UPGMA
- 4. Distance Methods Neighbor-Joining Algorithm
- 5. Maximum Parsimony and Maximum Likelihood
- 6. Bayesian Methods in Phylogenetics
- 7. Visualization, Analysis, and Evaluation of Obtained Trees
- 8. Monitoring Pathogen Evolution

Teaching methods

1. Lecture - multimedia presentation with examples presented on blackboard.

2. Laboratory:

- Performing practical tasks (including, raw data processing and analysis, tree construcion and visialisation, statistics, tree analysis and evaluation).

- Implemetation and validation of student's verions of phylogenetic algorithms.

- Preparing results in the form of reports.

- The use of popular software for phylogenetic analysis, combined with discussions on the functionality of selected analytical methods and algorithms used for constructing, visualizing, and evaluating phylogenetic trees

- Engaging in discussions

Bibliography

Basic:

Hall B., Łatwe drzewa filogenetyczne, WUW, Warszawa, 2008

Additional:

Felsenstein J., Inferring phylogenies, Sinauer Associates, , 2004 Warnow, T., Computational Phylogenetics: An Introduction to Designing Methods for Phylogeny Estimation. Cambridge, Cambridge University Press, 2017 Xia, X. A Mathematical Primer of Molecular Phylogenetics, Apple Academic Press Inc., 2020

Breakdown of average student's workload

	Hours	ECTS
Total workload	55	2,00
Classes requiring direct contact with the teacher	30	1,00
Student's own work (literature studies, preparation for laboratory classes/ tutorials, preparation for tests/exam, project preparation)	25	1,00