



## COURSE DESCRIPTION CARD - SYLLABUS

Course name

Populational Genetics

### Course

Field of study

Bioinformatics

Area of study (specialization)

Level of study

Second-cycle studies

Form of study

full-time

Year/Semester

1/1

Profile of study

general academic

Course offered in

Polish

Requirements

compulsory

### Number of hours

Lecture

30

Laboratory classes

30

Other (e.g. online)

Tutorials

Projects/seminars

### Number of credit points

5

### Lecturers

Responsible for the course/lecturer:

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Responsible for the course/lecturer:

### Prerequisites

The student starting this course should have a basic knowledge in basic computer programming. He/she should possess skills in solving basic problems that computer science deals with and the ability to acquire information from specific. They should also have basic knowledge of molecular biology, molecular genetics and statistics.

### Course objective

1. To provide students with basic knowledge in the field of population genomics, in particular: genetic and phenotypic variability and its organisation, influence of factors such as genetic drift, mutation, Darwinian selection.



2. To get students acquainted with bioinformatic tools, approaches and analyses used in population genomics research.
3. To get students acquainted with currently used methods of computational analysis in bioinformatics of nucleic acids on the example of RNA.

### Course-related learning outcomes

#### Knowledge

1. The student has a knowledge, with theoretical basis in bioinformatics in approaches, research and analyses concerning broadly defined population genomics and bioinformatics of nucleic acids on the example of RNA.
2. Knows the application of computer tools and programs in analysis of genetic data in population genomics and bioinformatics of nucleic acids.
3. Knows methods, techniques and tools used in the process of solving bioinformatics complex tasks, mainly of engineering character
4. Knows principles of planning research in bioinformatics field

#### Skills

1. Is able to retrieve and interpret the information from a variety of sources concerning scientific literature and the Internet.
2. Knows and is able identify, choose and apply in practice appropriate approaches and software for carrying out population genomics analysis. They will be able to interpret and discuss the results obtained by such analysis.
3. Is familiar with and able to apply in practice various methods dedicated to analysis, comparison, annotation and prediction of interactions in case of RNA molecules.
4. Is able to apply advanced techniques and computer tools to solve biological problems and evaluate their usefulness.
5. Is able to prepare a written report in Polish presenting the results of conducted analyses, as well as discuss them.
6. Formulates and tests hypotheses connected with bioinformatics problems.

#### Social competences

1. Is able to cooperate and to work in the group, taking different roles on in it.
2. Is able appropriately to determine priorities serving the performance of a task determined by oneself or other.
3. Is showing the creative posture in the working life and social.



## Methods for verifying learning outcomes and assessment criteria

Learning outcomes presented above are verified as follows:

Forming evaluation

a) in lectures verifying established effects of the education is being carried out through:

- filing the written test with 5-10 questions / tasks to solve - every task 0-4 pt (tasks can consist of a few subsections - there is a then set fragmentary score for every subsection).

b) in laboratories:

The final evaluation is determined on the average evaluations of the exercises and practical laboratories.

## Programme content

This course is devoted to a theoretical and practical introduction to the field of population genomics and to the bioinformatics of nucleic acids on the example of RNA. The following issues will be discussed during the lectures: historical background, importance of population genomics and its basic concepts; genetic and phenotypic variation (e.g. single and multiple inheritance, DNA polymorphism and its significance, molecular population genetics); organisation of genetic variation (e.g. Hardy-Weinberg's law, its extensions and testing methods, linkage disequilibrium and its causes); genetic drift (e.g. Wright-Fisher drift model, effective population size, founder and bottleneck effects, gene trees and coalescence); mutation and neutral theory (e.g. types of mutations, model of infinite number of alleles and sites, linkage to genetic drift and recombination); Darwinian selection (e.g. types of selection, selection in haploid and diploid organisms, link to mutation, equilibrium states); inbreeding and migration (e.g. divided populations, inbreeding and selective mating, types of migration and its models); molecular population genomics (including assessment of the rate of molecular sequence variation, types of nucleotide and amino acid substitutions, molecular clock, polymorphism, phylogeny, multigene families); evolutionary quantitative genetics and human genetics (e.g. genetic models, covariance, genetic correlation); basic methods for analysis and prediction of RNA secondary structure using existing bioinformatic and experimental approaches; methods of comparison, interaction and annotation of RNA gene sequences on the example of non-coding RNAs. Laboratories will consist of the analysis of biological and biomedical data, with a focus on getting to know programs, tools and approaches related to the topics discussed in the lecture and their applications.

## Teaching methods

Lecture illustrated by a multimedia presentation enriched with numerous examples

Laboratories: practical exercises in population genomics and dedicated RNA analysis, group work, discussion and problem analysis.

## Bibliography

Basic

Hartl D.I., Clark A.G., "Principles of Population Genetics", 2009



Additional

1. Avise J.C., "Molecular Markers, Natural History and Evolution", 2008
2. Baxevanis A.D., Ouellette B.F.F., "Bioinformatics: A Practical Guide to the Analysis of Genes and Proteins", 2005

**Breakdown of average student's workload**

	Hours	ECTS
Total workload	125	5,0
Classes requiring direct contact with the teacher	60	2,5
Student's own work (literature studies, preparation for laboratory classes/tutorials, preparation for tests/exam, project preparation) <sup>1</sup>	65	2,5

<sup>1</sup> delete or add other activities as appropriate