



OPIS PREDMETA / SUBJECT SPECIFICATION

Predmet: Subject Title:	HUMANA GENETIKA HUMAN GENETICS
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Študijski program Study programme	Študijska smer Study field	Letnik Year	Semester Semester
<i>Izobraževalna biologija</i> , pedagoški dvopredmetni študijski program 2. stopnje		1	Zimski
<i>Educational Biology</i> , pedagogical two stream study, 2 nd. degree		1	Winter

Univerzitetna koda predmeta / University subject code:

Predavanja Lectures	Seminar Seminar	Sem. vaje Tutorial	Lab. Vaje Lab. Work	Teren. vaje Field work	Samost. delo Individ. work	ECTS
15	15		15		45	3

Nosilec predmeta / Lecturer:

Jeziki / Predavanja / Lecture:
Languages: Vaje / Tutorial:

Pogoji za vključitev v delo oz. za opravljanje študijskih obveznosti:

Prerequisites:

Vsebina:

Contents (Syllabus outline):

2. Struktura, morfologija in klasifikacija humanih kromosomov.
3. Osnove citogenetike, molekularna citogenetika, klinična citogenetika – sindromologija.
4. Dedovanje, Mendelejevi zakoni, dominantno, recesivno dedovanje, atipični vzorci dedovanja.
5. Struktura in lastnosti DNA molekule.
6. DNA mutacije, popravljanje DNA napak.
7. Človeški genom, jedrni genom, genetski kod, zgradba gena, genske družine, tandemске ponovitve, mitohondrijski genom.
8. Genetska raznolikost med posamezniki in populacijami, polimorfizmi, genetsko neravnostezje.
9. Kompleksne bolezni, multifaktorsko dedovanje.
10. Somatske genetske bolezni: Citogenetika in molekularna genetika raka.
11. Izbrani primeri genetskih bolezni.
12. Genetsko svetovanje, prenatalna diagnostika, preimpantacijska genetika, genska terapija.
13. Genetsko testiranje v medicini.
14. Preprečevanje in zdravljenje genskih bolezni.
15. Etična vprašanja genetike v medicini.

2. Structure, morphology and classification of human chromosomes.
3. The principles of cytogenetics, molecular cytogenetics, clinical cytogenetics (syndromology).
4. Inheritance, Mendelian laws, dominant, recessive single-gene inheritance, atypical patterns of inheritance.
5. Structure and features of DNA molecule.
6. DNA mutations, repairing mechanisms.
7. Human genome, nuclear genome, genetic code, structure of the gene, gene families, tandem repeats, mitochondrial genome.
8. Genetic diversity between individuals and in populations, polymorphisms, linkage disequilibrium.
9. Complex diseases, multifactorial inheritance.
10. Somatic genetic diseases: Cytogenetics and molecular genetic of cancer.
11. Genetic diseases – selected cases.
12. Genetic counseling, prenatal diagnosis, preimplantation genetics, gene therapy.
13. Genetic testing in medicine.
14. Genetic disease prevention and treatment
15. Ethical issues in medical genetics.

Temeljni študijski viri / Textbooks:

1. Thompson & Thompson : Genetics in Medicine, W.B.Saunders Company., 6th ed. ISBN 0-7216-0244-4 and 7th ed. ISBN: 9781416030805, 2007.
2. Borut Peterlin, Karin Witzl: Humana genetika, Cankarjeva založba, 1 izdaja, 1 natis, ISBN 961-231-351-2, 2003.
3. Vogel, Motulsky, Human Genetics, Springer. 4th ed. ISBN 978-3-540-37653-8, 2009.

Cilji:

Predmet študenta seznanja z možnimi načini analize genoma in ugotavljanja genetskih napak. Študenta seznanja z zakonitostmi monogenskega dedovanja in boleznimi vezanimi z napakami posamičnih genov. Študenta seznanja z vlogo genetskih faktorjev pri vzrokih humanih boleznih ter prispevku k multifaktorskim boleznim, s kompleksno analizo dedovanja in prenosa genetske informacije ter dedovanjem. Poudarek predavanj je na aplikativni vlogi genetike v sodobnih medicinskih tehnikah, diagnostiki in genski terapiji.

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Predvideni študijski rezultati:

Znanje in razumevanje:

Pozna pravila in vrste monogenskega dedovanja.
 Pozna primere boleznih, ki so posledica sprememb v enem genu.
 Pozna normalni človeški kariotip, mehanizme kromosomskih napak, ki se pojavljajo v humanem genomu in posledične sindrome.
 Ve kaj je gen, kako nastajajo mutacije, kako se popravljajo in kako jih določamo.
 Pozna tehnike določanja kromosomskih in genskih sprememb, ki se uporabljajo v medicinski genetski diagnostiki.
 Razume, kaj je razlika med mutacijo in polimorfizmom ter pozna genetske variabilnosti med posamezniki in človeškimi populacijami.
 Pozna razloge za nastanek kompleksnih boleznih.
 Pozna osnovne mehanizme genetike raka.
 Pozna načine testiranja genetskih boleznih.
 Seznanja se z genetskim svetovanjem in načini preprečevanja genetskih boleznih ter gensko terapijo.
 Seznanja se z etičnimi problemi v medicinski genetiki.

Prenesljive/ključne spretnosti in drugi atributi:

1. Pozna, kako sestaviti kariogram po ISCN nomenklaturi.
2. Zna ločiti strukturno od numerične kromosomske spremembe.
3. Zna izolirati DNA molekulo iz periferne krvi in določiti njeno koncentracijo.
4. Pozna postopek verižne reakcije s polimerazo ter določitve produktov z gelsko elektroforezo.

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Objectives:

The courses offer to students the knowledge of major techniques used for genome analysis, and mutation analysis. The student is introduced to Mendelian laws, single-gene inheritance and diseases linked to monogenic disorders. The student is also introduced to the role of genetic factors in etiology of human hereditary disorders, the contribution to multifactorial diseases and understanding the complex information of functioning and transmission of genetic information. Special attention is given to diagnostic techniques, recombinant DNA technology and gene therapy.

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Intended learning outcomes:

Knowledge and Understanding:

- Knowledge of Mendelian laws, single-gene inheritance and diseases linked to monogenic disorders.
- Knowledge of human karyotype, mechanisms of chromosome rearrangements in human genome and the syndromes as their consequences.
- Understanding of the gene, mutations, repairing mechanisms and methods of detection.
- Knowledge of techniques used in medical genetic diagnostics.
- Understanding the differences between mutation and polymorphism and knowledge about genetic variations in individuals and human populations.
- Understanding of complex diseases.
- Understanding of principal mechanisms of cancer genetics.
- Knowledge of genetic tests.
- Understanding of genetic testing, preventing of genetic diseases and gene therapy.
- Understanding the ethical problems in medical genetics.

Transferable/Key Skills and other attributes:

1. Understanding of Mendelian laws, single-gene inheritance and diseases linked to monogenic disorders.
2. Understanding how to make karyotype according to ISCN nomenclature.
3. Knowledge of differences between structural and numerical chromosomal aberrations.
4. Knowledge of isolation of DNA molecules from peripheral blood and measuring its concentration.
5. Understanding of polymerase chain reaction

and determination of the product on gel electrophoresis.

Metode poučevanja in učenja:

- Predavanja: 33,3%
- Laboratorijske vaje : 33,3%
- Individualno delo: 33,3%

Learning and teaching methods:

- Lectures
- Laboratory excersises
- Individual work

Načini ocenjevanja:

- Ocena iz laboratorijskega dela in zaključni kolokvij.
- Ocena iz seminarske naloge.
- Pisni izpit

Delež (v %) /
Weight (in %)

10%
20%
70%

Assessment:

- Assessment of laboratory work
- Assessment of seminary work.
- Written exam

Materialni pogoji za izvedbo predmeta :

- Predavalnica
- Laboratorij

Material conditions for subject realization

- Lecture hall
- Laboratory

Obveznosti študentov:

(pisni, ustni izpit, naloge, projekti)

- Laboratorijsko delo
- Seminarsko delo
- Pisni izpit

Students' commitments:

(written, oral examination, coursework, projects):

- Laboratory work
- Seminar work
- Written exam