


INFORMAȚII PERSONALE

Andreescu Nicoleta Ioana

 Piata Eftimie Murgu, nr 2.Timisoara,RO-300041



EXPERIENȚA PROFESIONALĂ

Septembrie 2020-prezent

Conferentiar Universitar

Disciplina Genetica,Universitatea de Medicina si Farmacie „Victor Babes” Timisoara, Pta Eftimie Murgu Nr 2, Timisoara

sectorul de activitate

- Activitati didactice specifice

Ianuarie 2020-prezent

Conducator Doctorat

Universitatea de Medicina si Farmacie „Victor Babes” Timisoara, Pta Eftimie Murgu Nr 2, Timisoara

sectorul de activitate

- Activitati de cercetare specifice

Martie 2017-septembrie 2020

Sef de lucrari

Disciplina Genetica,Universitatea de Medicina si Farmacie „Victor Babes” Timisoara, Pta Eftimie Murgu Nr 2, Timisoara

sectorul de activitate

- Activitati didactice specifice

August 2018-prezent

Medic Primar Genetica Medicala

Centrul Regional de Genetica Medicala Timis, Spitalul Clinic Judetean de Urgenta pentru copii Timisoara

sectorul de activitate

- Activitate medicala, Laborator Genetica Medicala

Aprilie 2017-August 2018

Medic Specialist Genetica Medicala

Centrul Regional de Genetica Medicala Timis, Spitalul Clinic Judetean de Urgenta pentru copii Timisoara

sectorul de activitate

- Activitate medicala, Laborator Genetica Medicala

Aprilie 2014- Decembrie 2015

Cercetator postdoctorand

Disciplina Genetica,Universitatea de Medicina si Farmacie „Victor Babes” Timisoara, Pta Eftimie Murgu Nr 2, Timisoara

sectorul de activitate

- Activitati de cercetare specifice

Februarie 2013-2016

Medic Specialist Genetica Medicala, Coordonator Compartiment PCR

Spitalul Clinic Judetean de Urgenta Timisoara

sectorul de activitate

- Activitate medicala, Laborator Genetica Medicala

2011- 2017

Asistent universitar

Disciplina Genetica,Universitatea de Medicina si Farmacie „Victor Babes” Timisoara, Pta Eftimie Murgu Nr 2, Timisoara

sectorul de activitate

- Activitati didactice specifice

2007-2011

Preparator universitar

Disciplina Genetica,Universitatea de Medicina si Farmacie „Victor Babes” Timisoara, Pta Eftimie Murgu Nr 2, Timisoara

sectorul de activitate

- Activitati didactice specifice

- 2006-2010 **Doctorand**
Disciplina Genetica, Universitatea de Medicina si Farmacie „Victor Babes” Timisoara, Pta Eftimie Murgu Nr 2, Timisoara
sectorul de activitate
- Activitati de cercetare specifice

EDUCAȚIE ȘI FORMARE

- 2006-2010 **Doctor in Stiinte Medicale**
Disciplina Genetica, Universitatea de Medicina si Farmacie „Victor Babes” Timisoara, Pta Eftimie Murgu Nr 2, Timisoara
- Titlul tezei de doctorat: „Studiul prospectiv al depistarii afecțiunilor citogenetice prin metodologia diagnosticului antenatal”
- 24.02.2007-24.02.2012 **Medic Rezident**
Spitalul Clinic Judetean de Urgenta Timisoara
- Genetica medicala
- 01.01.2004 – 01.01.2005 **Medic stagiar**
Spitalul Clinica Municipal de Urgenta, Timisoara
- Medicina interna, oftalmologie
- 1997-2003 **Diploma de licenta**
Facultatea de Medicina, Universitatea de Medicina si Farmacie „Victor Babes” Timisoara
- Medicina Generala
- 1993-1997 **Diploma Bacalaureat**
Liceul „Traian Doda” Caransebes
- Profil Matematica Fizica

COMPETENTE PERSONALE

Limba maternă Limba romana

Alte limbi străine cunoscute

INTELEGERE		VORBIRE		SCRIERE
Ascultare	Citire	Participare la conversație	Discurs oral	
Limba engleza	B2	B2	B2	B2
Certificat emis de Universitatea de Vest Timisoara				
Limba Franceza	B2	B2	B2	B2
Certificat emis de Universitatea de Vest Timisoara				

Niveluri: A1/2: Utilizator elementar - B1/2: Utilizator independent - C1/2: Utilizator experimentat
Cadrul european comun de referință pentru limbi străine

Competențe de comunicare

- Spirit de echipa
- Capacitate de adaptare la noi conditii de lucru, capacitate de comunicare si interrelationare cu persoane din medii diferite

Competențe organizaționale/manageriale

- Initiativa si capacitate organizatorica

Competențe informatice

- Cunostinte de utilizare a calculatorului (Microsoft Office: Word/Excel/Power-Point)

INFORMATII SUPLIMENTARE

Membru al asociațiilor profesionale:

1. Societatea Română de Genetică
2. ECA
3. Societatea Europeană de Genetică(ESHG)
4. Societatea Romana de oftalmogenetica (membru fondator)

Specializări cursuri și calificări:

1. 2007, Oslo,Ullevål Hospital: Tehnici de cultura celulara, prelucrarea computerizata a cariotipului, tehnica FISH
2. 2008, Oslo, Radium Spital: Tehnici de cultura a celulelor tumorale, prelucrarea computerizata a cariotipului, tehnica FISH.
3. 2006-2007 Politehica Timisoara, Romania, Pedagogie
4. 15-20.05.2007,, Deficiențe de sexualizare”, UMFT Victor Babeș Timișoara
5. 16 noiembrie 2007“Tulburările legate de consumul de alcool,o problema de sanatate publica” desfășurat în Timisoara.
6. 13 noiembrie 2008 ”Coordonate de expunere ambientală la substanțe toxice” desfășurat la Timisoara.
7. mai 2008. UMF Timisoara Practical course of molecular cytogenetics – ISH,,Molecular Cytogenetics techniques using in situ hybridization”
8. 2009, UMFT, Real-time PCR-curs practic. Identificarea de biomarkeri cu aplicații în evaluarea predispoziției și orientarea tratamentului (polimorfismul ApoE și boala Alzheimer).
9. Septembrie 2010, Goldrain, South Tyrol, Italy. Clinical Cytogenetics
10. 2010, Institutul de Medicina Legală „Mina Minovici” București: Tehnici genetice moleculare MLPA, PCR, secvențiere genică.
11. 2011, Institutul Oncologic Cluj Napoca: Gene Microarray practical course. -16 septembrie 2014 Sesiune de instruire Managementul cercetării științifice
12. 18-19 noiembrie 2014 Sesiune de formare pentru dezvoltarea comunicării interpersonale, abilităților de comunicare, managementului timpului, leadershipului, dezvoltării personale
13. 27 februarie 2015 Sesiuni de formare on-line pe teme europene transversale
14. Iunie 2016 Curs Expert accesare fonduri structurale

Membru în bordul editorial:

1. Romanian Journal of Rare Diseases
2. Merit Research Journal of Medicine and Medical Sciences (MRJMMS)
3. Journal of Cancer Research and Treatment

Premii și distincții:

1. Premiul special Conferința Națională de Genetică Medicală, Sibiu, 24-26 septembrie 2009.
2. Premiul Cercetător Eminent 2015, Orizonturi Universitare Timișoara, decembrie 2015. Diploma de merit, Primăria Municipiului Timișoara, 2015.

Experiența acumulată (inclusiv experiența managerială) în programe/proiecte naționale/internaționale:

1. 2008-2011 membru al proiectului Correlation of clinical, genetics and epigenetics aspects for understanding the etiology Prader Willi/Angelman syndromes – a model of interdisciplinary approach of rare diseases in Romania- membru în proiect
2. Utilizarea modelelor nutri genomice pentru personalizarea tratamentelor dietetice în obezitate – NutriGen P_37- 684, cod SMIS 104852, 2016-2019 – Responsabil testare genetică
3. Implementarea transdisciplinară aplicativă a unor abordări moderne integrative ce corelează datele clinice, neurobiologice, farmacogenetice și neuroimagistice în vederea

dezvoltarii terapiilor personalizate in managementul tulburarilor psihotice – Responsabil proiect

4. Aplicarea si dezvoltarea unor abordari inovative, transdisciplinare de diagnostic si tratament in tulburarile de spectru autist, grant SC MedPsy DP RESEARCH SRL, contract nr 3F/29.11.2016, 2016-2019 – membru in proiect
5. Cresterea capacitatii de cercetare genetica si genomica in dezvoltarea perinatale si a copilului BM 29/2016 - membru in proiect
6. Science and Technology in childhood Obesity Policy, acronym: STOP, ID 774548
7. SARS-CoV-2 genome sequencing and phylogenetic analysis of circulating strains in Romania, P 2 - SP 2.1 - Soluții – 2020- 1 PN-III-P2-2.1-SOL-2020-0142

Lector la manifestari internationale,
nationale cu participare internationala,
nationale:

1. The 17th National Congress of Laboratory Medicine, 8-11 Septembrie 2010, Tg Jiu. Abordarea interdisciplinara a bolilor rare, 20 - 22 septembrie 2010.
2. International Conference for Human Reproductive Health, 26-28 Nov. 2010, Timisoara
3. World Kidney Day, 15 martie 2013, Timisoara
4. Al 52 lea Congres National de Cardiologie, 3-5 octombrie 2013, Sinaia
5. Conferința Națională de Management Medical Modern, Specializarea Genetică, 31 octombrie 2013, Bucuresti
6. Diagnosis and Discovery Workshop Partnership Opportunities with Central/Eastern Europe and the Middle East Prague, Czech Republic, December 3, 2013
7. 2014 Golden Helix Symposium Genomics of Rare Diseases October 30th - November 1st, 2014, Belgrade, Republic of Serbia
8. 2015- 11th Balkan Congress of Human Genetics, Belgrade, Serbia, Septembrie 2015. 2015 – 12th BANTAO Congress – DiaTransplant, 15-18 Octombrie, Opatija Croatia.

Organizare de evenimente științifice
(conferințe, workshop-uri):

1. Practical course of molecular cytogenetics – FISH „Molecular Cytogenetics techniques using in situ hybridization” 21st-23rd of May 2008.
2. The 3rd National Congress of Medical Genetics. 22-25 September 2010, Timisoara
3. A VII-a Conferinta Nationala de Genetica Medicala, 26-28 septembrie 2013, Paltinis
4. Al IV-a Congres National de Genetica Medicala, 25-27 septembrie 2014, Bucuresti
5. A VIII-A Conferinta Nationala de Genetica Medicala, 8-10 Octombrie 2015, Orastie.
6. A IX-A Conferinta Nationala de Genetica Medicala ALBA IULIA, 22-24 SEPTEMBER 2016
7. A X-A ConFERINTA DE Genetica Medicala cu participare internationala Craiova, 6-8 septembrie 2017
8. A XI A CONFERINȚĂ DE GENETICĂ MEDICALĂ CU PARTICIPARE INTERNAȚIONALĂ, TIMISOARA, 18-20 SEPTEMBRIE 2019
9. 14th Congress of the International Society of Nutrigenetics/Nutrigenomics 26-28 September 2021

Lista de lucrări

Titlul tezei de doctorat

Studiul prospectiv al depistării afecțiunilor citogenetice prin metodologia diagnosticului antenatal

Titlul tezei de abilitare

Genetica între cercetare, implicații clinice și noi provocări

Monografii, ghiduri:

- 1) Maria Puiu, Nicoleta Andreescu, Adela Chirita Emandi – editori: Ghid pentru situații de urgență în bolile rare ISBN 978-606-786-008-5, Editura Victor Babes Timisoara, 2016
- 2) Maria Puiu, Simona Farcas, Nicoleta Andreescu, Andreea Dobrescu, Diana Tiugan, Paul Tutac, Victoria Sacara, Natalia Usurelu, Iulia Coliban, Victoria Hlistun, Chiril Boiciuc, Dorif Alexandr, Turcan Doina, Daniela Blăniță, Natalia Barbova, Vladimir Egorov Abordarea bolilor genetice prin screening și diagnostic pre- și postnatal, Editura Victor Babes, Timisoara, 2019
- 3) Ruxandra Jurcut, Carmen Ginghina, Cardiomiopatii genetice, Editura Medicala Antaeus, 2018

Tratat internațional (autor capitol)

Andreescu N., Puiu M., Niculescu M. (2018) Effects of Dietary Nutrients on Epigenetic Changes in Cancer. In: Dumitrescu R., Verma M. (eds) Cancer Epigenetics for Precision Medicine. Methods in Molecular Biology, vol 1856. Humana Press, New York, NY

Tratat în editură recunoscută CNCSIS (membru colectiv autori)

Genetica Medicala ed 3 Mircea Covic, Dragos Stefanescu, Ionel Sandovici, Vlad Gorduză Editura: Polirom, 2017, ISBN 9789734665266

Cursuri universitare

- 1) De la fundamental la aplicativ în genetica stomatologică. Valerica Belengeanu, Florica Glavan, Dorina Stoicănescu, Noemi Meszaros, **Nicoleta Andreescu**, Cristina Bratu, Monica Stoian, Simona Farcas, Cristina Popa, Dragos Belengeanu Mihaita Opriteșcu. Editura Eurostampa, ISBN 978-973-687-644-8, Timisoara 2007.
- 2) Genetica Stomatologică - Baze teoretice și clinice. Valerica Belengeanu, Dorina Stoicănescu, Noemi Meszaros, **Nicoleta Andreescu**, Monica Stoian, Simona Farca, Cristina Bratu, Malina Popa, Cristina Popa, Dragoș Belengeanu, Stefania Dinu. Editura Eurostampa, ISBN 978-606-569-152-0, 2010.
- 3) Genetica anomaliilor dento-maxilo-faciale. Puiu Maria, Stoicanescu Dorina, Gug Cristina, Popa Cristina, Farcas Simona, **Andreescu Nicoleta**, Meszaros Noemi, Stoian Monica. Editura Victor Babes, Timisoara ISBN 978-606-8054-94-0, 2012.
- 4) Genetics in clinical dentistry - Stoicanescu Dorina, **Andreescu Nicoleta**, Editura Eurostampa, 2013, ISBN 978-606-569-666-2.
- 5) Curs de Genetică Medicală. Maria Puiu, Dorina Stoicanescu, Cristina Gug, Simona Farcas, Cristina Popa, **Nicoleta Andreescu**, Adela Chirita-Emandi, Andreea Dobrescu. Editura Eurostampa, 2016, ISBN 978-606-32-0296-4.
- 6) Curs de GENETICĂ pentru Asistență Medicală Generală MARIA PUIU, DORINA STOICĂNESCU, CRISTINA GUG, SIMONA FARCAȘ, CRISTINA POPA, **NICOLETA ANDREESCU**, ADELA CHIRIȚĂ-EMANDI, ANDREEA DOBRESCU, Editura “Victor Babeș” Timișoara, 2017, ISBN 978-606-786-043-6
- 7) Genetics in Clinical Dentistry. 2nd Edition Stoicanescu Dorina, **Andreescu Nicoleta**, Eurostampa, 2019, ISBN 978-606-32-0773-0

Suport de studiu pentru lucrări practice

- 1) Aplicații practice în genetica medicală, ed.I Valerica Belengeanu, Maria Puiu, Dorina Stoicănescu, Cristina Gug, Mirela Mihaescu, Simona Farcaș, Cristina Popa Monica Stoian, **Nicoleta Andreescu**, Noemi Meszaros. Editura Eurostampa, ISBN 978-973-687-676-9. Timisoara 2008.
- 2) Aplicații practice în genetica medicală, ed.II revizuită Valerica Belengeanu, Maria Puiu, Dorina Stoicănescu, Cristina Gug, Mirela Mihaescu, Simona Farcaș, Cristina Popa Monica Stoian, **Nicoleta Andreescu**, Noemi Meszaros. Editura Eurostampa, ISBN 978-973-687-762-9. Timisoara 2008.
- 3) Aplicații practice în genetica medicală, ed.III revizuită Valerica Belengeanu, Maria Puiu, Dorina Stoicănescu, Cristina Gug, Mirela Mihaescu, Simona Farcaș, Cristina Popa Monica Stoian, **Nicoleta Andreescu**, Noemi Meszaros Editura Eurostampa, ISBN 978-973-687-915-9. Timisoara 2009.
- 4) Genetica medicală în practica clinică. Valerica Belengeanu, Maria Puiu, Dorina Stoicănescu, Cristina Gug, Simona Farcaș, Cristina Popa Monica Stoian, **Nicoleta Andreescu**, Noemi Meszaros. Eurostampa Publishing, ISBN 978-606-569-213-8. Timisoara 2011.

- 5) Genetica medicala – caiet lucrari practice. Puiu Maria, Stoicanescu Dorina, Gug Cristina, Popa Cristina, Farcas Simona, **Andreescu Nicoleta**, Adela Emandi-Chirita, Corina Pienar, Meszaros Noemi. Editura Eurostampa, ISBN 978-606-569-563-4, 2013
- 6) Aplicații practice de GENETICĂ pentru Asistență Medicală Generală MARIA PUIU, DORINA STOICĂNESCU, CRISTINA GUG, SIMONA FARCAȘ, CRISTINA POPA, **NICOLETA ANDREESCU**, ADELA CHIRIȚĂ-EMANDI, ANDREEA DOBRESCU, Editura “Victor Babeș” Timișoara, 2017, ISBN 978-606-786-044-3
- 7) Genetică - Aplicații practice Maria Puiu, Dorina Stoicanescu, Cristina Gug, Simona Farcas, Cristina Popa, **Nicoleta Andreescu**, Adela Chirita-Emandi, Andreea Dobrescu, Alexandra Mihăilescu. Editura Eurostampa, 2019 ISBN 978-606-32-0670-2

Lucrari ISI autor principal

- 1) D. Belengeanu, Cristina Bratu, Monica Stoian, A. Motoc, Eli Ormerod, Angela Codruța Podariu, Simona Farcas, **Nicoleta Andreescu** The heterogeneity of craniofacial morphology in Prader-Willi patients. Rom J Morphol Embryol 2012, 53(3):527–532. ISSN 1220-0522 **IF=0.620**
- 2) SIMONA FARCAȘ, C.D. CRIȘAN, **NICOLETA ANDREESCU***, MONICA STOIAN, A. G. M. MOTOC (***autor corespondenta**) Structural chromosomal anomalies detected by prenatal genetic diagnosis: our experience, Rom J Morphol Embryol 2013, 54(2):377–383 ISSN 1220-0522 **IF= 0.723**
- 3) DORINA STOICĂNESCU, **NICOLETA ANDREESCU***, ALINA BELENGEANU, NOEMI MESZAROS, MĂRIOARA CORNIANU Rom (***autor corespondenta**) Assessment of p53 and HER-2/neu genes status and protein products in oral squamous cell carcinomas Rom J Morphol Embryol 2013, 54(4):1107–1113 ISSN 1220-0522 **IF= 0.723**
- 4) Laura Alexandra Nussbaum, **Nicoleta Andreescu***, Liliana Nussbaum, Raluca Gradinaru, Maria Puiu (***autor corespondenta**) Ethical Issues Related to Early Intervention in Children and Adolescents with Ultra High Risk for Psychosis: Clinical Implications and Future Perspectives Rev Rom Bioet, 2014, vol. 12(3), 64-81 ISSN 1583-5170 **IF=0.462**
- 5) Laura Nussbaum, Raluca Gradinaru, **Nicoleta Andreescu***, Victor Dumitrascu, Anca Tudor, Liana Suci, Radu Stefanescu, Maria Puiu (***autor corespondenta**) The Response to Atypical Antipsychotic Drugs in Correlation with the CYP2D6 Genotype: Clinical Implications and Perspectives Farmacia, 2014, vol. 62(6),1191-1201 ISSN 0014-8237 **IF=1.005**
- 6) Razvan Vladimir Socolov, **Nicoleta Ioana Andreescu**, Ana Maria Haliciu, Eusebiu Vlad Gorduza, Florentin Dumitrache, Raluca Anca Balan, Maria Puiu, Mihaela Amelia Dobrescu, Demetra Gabriela Socolov (*** autor de corespondenta**) Intrapartum diagnostic of Roberts syndrome – case presentation Rom J Morphol Embryol 2015, 56(2):585–588 ISSN 1220-0522 **IF 0.811**
- 7) **Nicoleta Andreescu**, Laura Nussbaum, Lavinia Maria Hoge, Raluca Gradinaru, Calin Muntean, Radu Ștefanescu, Maria Puiu Antipsychotic treatment emergent adverse events in correlation with the pharmacogenetic testing and drug interactions in children and adolescents with schizophrenia and bipolar disorder FARMACIA, 2016, 64(5):736-744 ISSN 0014-8237 **IF= 1.348**
- 8) **Nicoleta Ioana Andreescu**, Mirela Cosma, Simona Sorina Farcas, Monica Stoian, Daniela-Georgiana Amzar, Maria Puiu Assessment of chromosomal aneuploidies in sperm of infertile males by using FISH technique Rom J Morphol Embryol 2016, 57(1):173–178 ISSN 1220-0522 **IF=0.670**
- 9) Raluca Claudia Gradinaru, **Nicoleta Ioana Andreescu***, Laura Alexandra Nussbaum, Simona Sorina Farcas, Victor Dumitrascu, Liana Suci, Maria Puiu (***autor de corespondenta**), -759C/T polymorphism of the HTR2C gene is not correlated with atypical antipsychotics-induced weight gain, among Romanian psychotic patients Rom J Morphol Embryol 2016, 57(4):1–7 ISSN 1220-0522 **IF=0.670**
- 10) F Stoica, D Ionescu, A Heghes, C Trandafirescu, **N Andreescu***, A Tudor, Sebastian Olariu, Alina Stanciu, Mihaela Galea, Maria Puiu (***autor de corespondenta**) Vascular Endothelial Growth Factor Gene Polymorphism-Susceptibility Predictor for Severe Retinopathy of Prematurity? REVISTA DE CHIMIE 2016, 67 (12), 2522-2525 ISSN 2537-5733 **IF=1.232**
- 11) Dragoș Erdelean, Simona Sorina Farcas, Vladimir Poroș, **Nicoleta Ioana Andreescu***, Izabella Erdelean, Andreea Iulia Dobrescu, Laura Alexandra Nussbaum, Lavinia Maria Hoge, Dan Navolan, Paul Tutac, Maria Puiu (***autor corespondenta**) Association between thrombophilia gene polymorphisms and recurrent pregnancy REV.CHIM., 2018, 69(11):3122-3125. ISSN 2537-5733 **IF=1.605**
- 12) Adela Chirita Emandi, Diana Munteanu, **Nicoleta Andreescu***, Paul Tutac, Corina Paul, Iulian Puiu Velea, Agneta Maria Pusztai, Victoria Hlistun, Chiril Boiciuc, Victoria Sacara, Lorina Vudu, Natalia Usurelu, Maria Puiu (***autor corespondenta**) No clinical utility of common polymorphisms in IGF1, IRS1, GCKR, PPARG, GCK1 and KCTD1 genes previously associated with insulin resistance in overweight children from Romania and Moldova Journal of Pediatric Endocrinology and Metabolism, 2018, <https://doi.org/10.1515/jpem-2018-0288> **IF=1.239**
- 13) Vlad Serafim, Diana-Andreea Tiugan, **Nicoleta Andreescu***, Alexandra Mihăilescu, Corina Paul, Iulian Velea, Maria Puiu, Mihai Dinu Niculescu (***autor corespondenta**) Development and Validation of a LC-MS/MS-Based Assay for Quantification of Free and Total Omega 3 and 6 Fatty Acids from Human Plasma Molecules 2019, 24, 360; doi:10.3390/molecules24020360 **IF=3.06**

- 14) Grădinaru R*, **Andreescu N***, Nussbaum L, Suci L, Puiu M (* **equal contribution**) Impact of the CYP2D6 phenotype on hyperprolactinemia development as an adverse event of treatment with atypical antipsychotic agents in pediatric patients. *Ir J Med Sci.* 2019 Feb 15. Doi: 10.1007/s11845-019-01985-x. **IF=1.031**
- 15) NOEMI MESZAROS, **NICOLETA IOANA ANDREESCU***, SIMONA SORINA FARCAS, ANDREEA IULIA DOBRESU, LAVINIA ELENA STELEA, ENDRE MATHE, ANCA PORUMB, MARIA PUIU (***autor corespondenta**) TERT Genotyping for Evaluation of Reproduction Failure REV. CHIM.(Bucharest 2019, 70(1): 195-198 **IF=1.412**
- 16) Farcas, Simona; **Andreescu, Nicoleta***; Amzar, Daniela; et al. (*autor corespondenta) Cytogenetic Study of Spontaneous Abortions in the Western part of Romania REVISTA DE CHIMIE, 2019, 70(3): 1000-1004. **IF=1.605**
- 17) Serafim V, Chirita-Emandi A, **Andreescu N***, Tiugan DA, Tutac P, Paul C, Velea I, Mihailescu A, Șerban CL, Zimbru CG, Puiu M, Niculescu MD. (***autor corespondenta**) Single Nucleotide Polymorphisms in *PEMT* and *MTHFR* Genes are Associated with Omega 3 and 6 Fatty Acid Levels in the Red Blood Cells of Children with Obesity. *Nutrients.* 2019 Oct 30;11(11). If= **4.17**
- 18) Chirita-Emandi A*, **Andreescu N***, Zimbru CG, Tutac P, Arghirescu S, Serban M, Puiu M. (* **equal contribution**) Challenges in reporting pathogenic/potentially pathogenic variants in 94 cancer predisposing genes - in pediatric patients screened with NGS panels. *Sci Rep.* 2020 Jan 14;10(1):223. doi: 10.1038/s41598-019-57080-9. **IF=4.011**
- 19) Chirita-Emandi A, Serban CL, Paul C, **Andreescu N***, Velea I, Mihailescu A, Serafim V, Tiugan DA, Tutac P, Zimbru C, Puiu M, Niculescu MD (***autor corespondenta**) CHDH-PNPLA3 Gene-Gene Interactions Predict Insulin Resistance in Children with Obesity. *Diabetes Metab Syndr Obes.* 2020, 19(13):4483-4494 doi 10.2147/DMSO.S277268 **IF=3.168**
- 20) Alin Viorel Istodor, Laura-Cristina Rusu, Gratiela Georgiana Noja, Alexandra Roi, Ciprian Roi, Emanuel Bratu, Georgiana Moise, Maria Puiu, Simona Sorina Farcas, **Nicoleta Ioana Andreescu*** An observational study on cephalometric characteristics and patterns associated with the Prader–Willi Syndrome: A structural equation modelling and network approach *Applied Sciences* 2021, 11 (7) 3177. doi 10.3390/app11073177 **IF=2.679**
- 21) Mihailescu, A.; Serafim, V.; Paul, C.; **Andreescu, N.**; Tiugan, D.-A.; Tutac, P.; Velea, I.; Zimbru, C.G.; Serban, C.L.; Ion, A.I.; David, V.L.; Ionescu, A.; Puiu, M.; Niculescu, M.D. Docosahexaenoic Acid and Eicosapentaenoic Acid Intakes Modulate the Association of *FADS2* Gene Polymorphism rs526126 with Plasma Free Docosahexaenoic Acid Levels in Overweight Children. *Appl. Sci.* 2021, 11, 9845. <https://doi.org/10.3390/app11219845> **IF=2.679**
- 22) Roi, A.; **Andreescu, N.I.**; Roi, C.I.; Negrut, iu, M.-L.; Sinescu, C.; Ravis, , M.; Boruga, M.V.; Rusu, L.-C. Comparative Analysis of COL9A1 Genotyping in Oral Squamous Cell Carcinoma Diagnosis: A Pilot Study. *Appl. Sci.* 2021, 11, 11102. <https://doi.org/10.3390/app112311102> **IF=2.679**

Lucrari ISI coautor

- 1) Valerica Belengeanu, Dorina Stoicanescu, **Nicoleta Andreescu**, Maria Constantinescu, Anca Muresan. Syndromic 46,XY disorder of sexual development. *Acta endocrinologica* Vol IV, No. 1, January-March, 2008. ISSN 1841-0987. **IF=0.052**
- 2) Valerica Belengeanu, Dorina Stoicanescu, Monica Stoian, **Nicoleta Andreescu**, Camelia Budisan. Ichthyosis congenital, harlequin fetus type: a case report *Advances in medical Science*, vol 54(1), 2009, pp113-115. **IF=0.798**
- 3) V. Belengeanu, H. Viskari, J. Tallila, J. Lahtela, S. Farcas, **N. Andreescu**, M. Stoian, C.L. Bohiltea, and J.P. Fryns. Lethal evolution of a newborn with consistent Features of hydrolethrus syndrome –Romanian patient. *Genetic Counseling*, Vol. 22, No 3, 2011, pp 293-304. **IF=0.505**
- 4) Dorina Stoicanescu, Valerica Belengeanu, Monica Stoian, **Nicoleta Andreescu**, Edward Seclaman, Andrei Anghel, Alina Belengeanu Clinical value of molecular testing in patients with Wilson disease Valoarea clinică a testelor moleculare la pacienții cu boala Wilson *Revista Română de Medicină de Laborator*, 2011,19(4):373-379 **IF=0.091**
- 5) Belengeanu V, Gamage TH, Farcas S, Stoian M, **Andreescu N**, Belengeanu A, Frengen E, Misceo D. A de novo 2.3Mb deletion in 2q24.2q24.3 in a 20-month-old developmentally delayed girl. *Gene* 2014 Apr 10;539(1):168-72 **IF=2.138**
- 6) Laura Alexandra Nussbaum, Victor Dumitrascu, Anca Tudor, Raluca Grădinaru, **Nicoleta Andreescu**, Maria Puiu Molecular Study of Weight Gain Related to Atypical Antipsychotics: Clinical Implications of the CYP2D6 Genotype *Rom J Morphol Embryol*, 2014, vol. 55(3), 877-884 ISSN 1220-0522 **IF=0.659**
- 7) Dumache R, Rogobete AF, **Andreescu N**, Puiu M. Genetic and Epigenetic Biomarkers of Molecular Alterations in Oral Carcinogenesis. *Clin Lab.* 2015;61(10):1373-81 **IF=0.936**
- 8) Laura Nussbaum, **Nicoleta Andreescu**, Lavinia Maria Hoge, Călin Muntean, Radu Ștefănescu, Maria Puiu Pharmacological and Clinical Aspects of Efficacy, Safety and Tolerability of Atypical Antipsychotic Medication in Child and Adolescent Patients with Schizophrenia and Bipolar Disorders *Farmacia*, 2016, vol. 64(6), 868-875 ISSN 0014-8237 **IF=1.348**
- 9) Laura Alexandra Nussbaum, Lavinia Maria Hoge, **Nicoleta Ioana Andreescu**, Raluca Claudia Grădinaru, Maria Puiu, Andrei Todica The Prognostic and Clinical Significance of Neuroimaging and Neurobiological Vulnerability Markers in Correlation with the Molecular Pharmacogenetic Testing in Psychoses and Ultra High-Risk Categories *Rom J Morphol Embryol*, 2016, vol. 57(3), 959-967 ISSN 1220-0522 **IF=0.670**
- 10) Hoge LM, Nussbaum LA, Chiriac DV, Ageu LȘ, **Andreescu NI**, Grigoraș ML, Folescu R, Bredicean AC, Puiu M, Roșca ECI, Simu MA, Levai CM. Integrative clinico-biological, pharmacogenetic, neuroimaging, neuroendocrinological and psychological correlations in depressive and anxiety disorders. *Rom J Morphol Embryol.* 2017;58(3):767-775. **IF=0.912**

- 11) LAURA NUSSBAUM, LAVINIA MARIA HOGEA, DANIELA CĂLINA, **NICOLETA ANDREESCU**, RALUCA GRĂDINARU, RADU ȘTEFĂNESCU, MARIA PUIU MODERN TREATMENT APPROACHES IN PSYCHOSES. PHARMACOGENETIC, NEUROIMAGISTIC AND CLINICAL IMPLICATIONS FARMACIA, 2017, 65(1):75-81. **IF=1.507**
- 12) Serafim V, Shah A, Puiu M, **Andreescu N**, Coricovac D, Nosyrev A, Spandidos DA, Tsatsakis AM, Dehelean C, Pinzaru. Classification of cancer cell lines using matrix-assisted laser desorption/ionization time-of-flight mass spectrometry and statistical analysis. Int J Mol Med. 2017 Oct;40(4):1096-1104. **IF=2.784**
- 13) DAN NAVOLAN, CRINGU ANTONIU IONESCU, ADRIAN CARABINEANU, FLORIN BIRSAȘTEANU, OCTAVIAN CRETU, FLORIN SZASZ, SIMONA VLADAREANU, IOANA CIOHAT, RAMONA GIDEA, DRAGOS NEMESCU, SIMONA FARCAS, **NICOLETA ANDREESCU**, SEBASTIAN SIMU, DANA STOIAN Influence of Weight of Pregnant Women on First Trimester Biochemical Markers Values REV.CHIM.(Bucharest), 2017,68(12)2836-2838. **IF=1.412**
- 14) Florina Stoica, Adela Chirita-Emandi, **Nicoleta Andreescu**, Alina Stanciu, Cristian G. Zimbru, Maria Puiu Clinical relevance of retinal structure in children with laser-treated retinopathy of prematurity versus controls – using optical coherence tomography Acta Ophthalmol, 2018: 96: e222–e228, doi: 10.1111/aos.13536. **IF=3.153**
- 15) LUMINIȚA ȘTEFANIA AGEU, CODRINA MIHAELA LEVAI, **NICOLETA IOANA ANDREESCU**, MIRELA LOREDANA GRIGORAȘ, LAVINIA MARIA HOGEA, DANIELA VERONICA CHIRIAC, ROXANA FOLESCU, ANA CRISTINA BREDICEAN, LILIANA MARIA NUSSBAUM, VIRGIL RADU ENĂTESCU, VLADIMIR POROCH, VIOREL LUPU, MARIA PUIU, LAURA ALEXANDRA NUSSBAUM Modern molecular study of weight gain related to antidepressant treatment: clinical implications of the pharmacogenetic testing Rom J Morphol Embryol 2018, 59(1):165–173 **IF=1.5**
- 16) FLORIN SZASZ, CODRINA LEVAI, DAN NAVOLAN, SIMONA FARCAS, **NICOLETA ANDREESCU**, FLORIN BIRSAȘTEANU, CLAUDIA MEHEDINTU*, CRINGU ANTONIU IONESCU, ROXANA BOHILTEA, ADRIAN CARABINEANU, DRAGOS NEMESCU, SEBASTIAN SIMU, DANA STOIAN Weight of Pregnant Women and their Influence on Second Trimester Biochemical Markers REV.CHIM.(Bucharest), 2018, 69(2):529-532. **IF=1.605**
- 17) Cristian Jinca, Carmen Angela Maria Petrescu, Estera Boeriu, Andrada Oprisoni, Loredana Balint-Gib, Mihaela Baica, Cristina Popa, **Nicoleta Andreescu**, Margit Serban, Emilia Ursu, Smaranda Arghirescu The impact of immunological and biomolecular investigations on the outcome of children with acute lymphoblastic leukemia - experience of IIIrd Paediatric Clinic Timisoara Revista Română de Medicină de Laborator, 2018, 26(1)77-85 **IF=0.8**
- 18) Simona Farcas, Dragos Erdelean, Flavia Anne-Elise Szekely, Dan Navolan, **Nicoleta Andreescu**, Andreea Cioca A rare case of partial trisomy 8q24.12-q24.3 and partial monosomy of 8q24.3: Prenatal diagnosis and clinical findings Taiwanese Journal of Obstetrics & Gynecology 58 (2019) 36-39 **IF=1.2**
- 19) IULIUS JUGANARU, CONSTANTIN TUDOR LUCA, ANDREEA-IULIA DOBRESCU, OANA VOINESCU, MARIA PUIU, SIMONA FARCAS, **NICOLETA ANDREESCU**, MIRCEA IURCIUC A Non-invasive, Easy to Use Medical Device for Arterial Stiffness REV.CHIM.(Bucharest), 2019, 70(2):642-645. **IF=1.605**
- 20) Borcan F, Chirita-Emandi A, Andreescu NI, Borcan LC, Albulescu RC, Puiu M, Tomescu MC. Synthesis and preliminary characterization of polyurethane nanoparticles with ginger extract as a possible cardiovascular protector. Int J Nanomedicine. 2019 May 21;14:3691-3703. doi: 10.2147/IJN.S202049. **IF=4.471**
- 21) Alexandra Roi, Ciprian Ioan Roi, **Nicoleta Ioana Andreescu**, Mircea Riviș, Ioana Daniela Badea, Noemi Meszaros, Laura Cristina Rusu, Stela Iurciuc Oral cancer histopathological subtypes in association with risk factors: a 5-year retrospective study Rom J Morphol Embryol. 2020 Oct-Dec;61(4):1213-1220. doi: 10.47162/RJME.61.4.22. **IF=1.033**
- 22) Adriana Cojocaru, Lavinia Maria Hogeia, Vladimir Porocho, Mihaela Adriana Simu, Virgil Radu Enatescu, Roxana Jeleriu, **Nicoleta Ioana Andreescu**, Maria Puiu, Bogdan Gheorghe Hogeia, Mirela Grigoras, Roxana Folescu, Carmen Lăcrămioara Zamfir, Ileana Enatescu and Laura Alexandra Nussbaum Effectiveness of Psychostimulant and Non-Psychostimulant Drug Therapy in the Attention Deficit Hyperactivity Disorder Appl. Sci. 2021, 11, 502. Doi: 10.3390/app11020502 **IF=2.679**
- 23) Adela Chirita-Emandi, **Nicoleta Andreescu**, Cristina Popa, Alexandra Mihailescu, Anca-Lelia Riza, Razvan Plesea, Mihai Ioana, Smaranda Arghirescu, Maria Puiu Biallelic variants in *BRCA1* gene cause a recognisable phenotype within chromosomal instability syndromes reframed as BRCA1 deficiency Journal of Medical Genetics 2021; **58**:648-652. Doi 0.1136/jmedgenet-2020-107198 **IF=6.318**
- 24) Enache, A.; Ciocan, V.; Muresan, C.O.; Cut, T.G.; Novacescu, D.; Paul, C.; **Andreescu, N.**; Mihailescu, A.; Raica, M.; Dumache, R. Postmortem Documentation of SARS-CoV-2 in Utero and Postpartum Transmission, through Amniotic Fluid, Placental, and Pulmonary Tissue RT-PCR. Appl. Sci. 2021, 11, 9505. doi: 10.3390/app11209505 **IF=2.679**
- 25) MARIA CLAUDIA JURCĂ1,2), OANA ALEXANDRA IUHAS2), MARIA PUIU3,4), ADELA CHIRIȚĂ-EMANDI3,4), **NICOLETA IOANA ANDREESCU**3,4), CODRUȚA DIANA PETCHEȘI 1), ALEXANDRU DANIEL JURCĂ1), IOAN MAGYAR1), SÂNZIANA IULIA JURCĂ5), KINGA KOZMA1,2), EMILIA MARIA SEVERIN6), MARIUS BEMBEA2) Cardiofaciocutaneous syndrome – a longitudinal study of a case over 33 years: case report and review of the literature Rom J Morphol Embryol 2021, 62(2):in press, doi: 10.47162/RJME.62.2.y **IF=1.033**

- 1) Nicoleta Andreescu, Dorina Stoicănescu, Alina Belengeanu, Simona Farcaș, Cristina Popa, Monica Stoian, Valerica Belengeanu. Unbalanced karyotype in a human foetus due to a recurrent familial translocation. *Analele Universității din Oradea - Fascicula Biologie*, Tom. XVII / 1, 2010, pp. 9-13.
- 2) Nicoleta Andreescu, Valerica Belengeanu, Simona Farcaș, Monica Stoian, I. Cioata, Miruna Munteanu, Elena Bernad. Rapid prenatal diagnosis using fish on uncultured amniotic fluid cells and chorionic villus sampling – experience on 60 cases. *Acta Medica Marisiensis*. Volume 56 Number 3, 2010, pp 243-245. ISSN. 2068-3324.
- 3) Monica Stoian, Valerica Belengeanu, Marioara Boia, Nicoleta Andreescu, Simona Farcaș, The role of fluorescence in situ hybridization in assessing the cytogenetically diagnosis in cryptical mosaicism aneuploidies, *Jurnalul Pediatriei*, Year X, Vol. X, Nr. 39-49, July-December 2007, pg 6-11.
- 4) Simona Farcaș, Valerica Belengeanu, Monica Stoian, Dorina Stoicanescu, Cristina Popa, Nicoleta Andreescu. CONSIDERATIONS REGARDING THE IMPLICATION OF POLYMORPHIC VARIANTS AND CHROMOSOMAL INVERSIONS IN RECURRENT MISCARRIAGE *Jurnalul Pediatriei*, Year X, Vol. X, Nr. 37-38, January-June 2007, pg 6-10.
- 5) Eli Ormerod, Valerica Belengeanu, Monica Stoian, Nicoleta Andreescu, Simona Farcaș, Cristina Popa, Mariana Banateanu, Alina Belengeanu. Nijmegen breakage syndrome –clinico-cytogenetic pattern. *Jurnalul Pediatriei*, Year XII, Vol. XII, Nr. 45-46, January-June 2009, pg 19-24.
- 6) Monica Stoian, Valerica Belengeanu, Simona Farcaș, Nicoleta Andreescu, Cristina Popa, Marioara Boia, Mihaita Opritescu, Eli Ormerod. Lejeune syndrome-a microdeletion syndrome-case report. *Jurnalul Pediatriei*, vol. XII, Nr. 47-48, July-December 2009, pg 16-19.
- 7) Noemi Meszaros, Dragoș Belengeanu, Dorina Stoicănescu, Nicoleta Andreescu, Simona Farcaș, Monica Stoian, Mariana Cevei. Analyses of numerical aberrations of chromosome 17 and TP53 gene deletion/amplification in human oral squamous cell carcinoma using dual-color fluorescence in situ hybridization *Analele Universitatii din Oradea, Fascicula de Biologie TOM XVII/1*, 2010, pag. 142-146.
- 8) Valerica Belengeanu, Marioara Boia, Simona Farcaș, Cristina Popa, Monica Stoian, Alina Belengeanu, Nicoleta Andreescu, Philippe Vago, Carole Goumy. Trisomy 8 mosaicism with atypical phenotypic features. *Jurnalul Pediatriei – Year XIII*, Vol. XIII, Nr. 51-52, july-december 2010, pg. 36-39.
- 9) Monica Stoian, Valerica Belengeanu, Maria Puiu, Natalia Cucu, Simona Farcaș, Nicoleta Andreescu, D. Belegeanu. Clinical and genetic investigations of 20 patients evaluated for Prader-Willi syndrome. *Acta Medica Marisiensis*. Volume 56, Number 2, 2010 pp. 69-72. ISSN. 2068-3324.
- 10) Cristina A. Popa, Hortensia Ioniță, Nicoleta Andreescu, Alina Belengeanu. Philadelphia chromosome in acute leukemia. *Medicine in evolution*. Volum XV, Nr. 2/2010 ISSN 2065-376X, pp 89-93.
- 11) Noémi Mészáros, Belengeanu Alina, Lazăr Elena, Cornianu Mărioara, Stoicănescu Dorina, Andreescu Nicoleta, Farcaș Simona, Stoian Monica, Popa Cristina. Correlation between expression of p53 mutant nuclear phosphoprotein, gene deletion and histopathological features in oral squamous cell carcinoma. *Acta Medica Marisiensis*. Volume 56 Number 4, 2010, pp 304-307. ISSN 2068-3324.
- 12) Catalina Giurgi-Onocu, Nicoleta Andreescu*, Cristina Bredicean, Doriană Misceș, Alina Belengeanu The schizophrenia pattern: associations with the shank3 gene – a case presentation. *Medicine in Evolution* Volume XVII, No. 4, 2012. (*autor corespondentă)
- 13) Simona Farcaș, Valerica Belengeanu, Monica Stoian, Nicoleta Andreescu*, Dragoș Belengeanu, Marioara Boia. Variable prognosis in trisomy 18 (Edwards Syndrome) -3 clinical cases presentation *Jurnalul Pediatriei*-Year XVI, Vol. XVI, Nr. 61-62, january-june 2013. (*autor corespondentă)
- 14) Raluca Grădinaru, Nicoleta Andreescu, Laura Nussbaum, Simona Farcaș, Maria Puiu Pharmacogenetic aspects which influence the pharmacokinetic properties of atypical antipsychotics – preliminary study. *Jurnalul Pediatriei*, Year XVI, Vol. XVI, Nr. 64, october-december 2013
- 15) Ramona Albușescu, Florina Stoica, Nicoleta Andreescu, Maria Puiu Retinopathy of prematurity – risk factors for evolution. *Jurnalul Pediatriei*, Year XVI, Vol. XVI, Nr. 64, october-december 2013
- 16) Daniela NEDELCU, Nicoleta ANDREESCU*, Estera BOERIU, Radu STEFANESCU, Smaranda ARGHIRESCU, Maria PUIU. Retrospective Study on Osteosarcoma and Ewing Sarcoma – Our Experience, *MAEDICA – a Journal of Clinical Medicine*, 2014; 9(2): 151-156 (*autor corespondentă)
- 17) MIRELA COSMA, SIMONA FARCAS, NICOLETA ANDREESCU*, CIPRIAN DORU CRISAN, OCTAVIA CIONCA, MARIA PUIU CORRELATIONS BETWEEN HETEROMORPHIC CHROMOSOMAL VARIANTS AND INFERTILITY *Medicine in Evolution* Volume XX, No. 2, 2014. (*autor corespondentă).
- 18) MIRELA COSMA, SIMONA FARCAS, MONICA STOIAN, DANIELA AMZAR, NICOLETA ANDREESCU*, MARIA PUIU MALE INFERTILITY-CYTOGENETIC FINDINGS IN A COHORT OF PATIENTS FROM WESTERN PART OF ROMANIA *Medicine in Evolution* Volume XX, No. 3, 2014 (*autor corespondentă).

- 19) Florina STOICA; Corina LADARIU; Marie-Jeanne KOOS; Alina STANCIU; Gabriela OLARIU; Nicoleta ANDREESCU; Maria PUIU Refractive and Visual Outcome after Laser-Treated Retinopathy of Prematurity in Western Romania MAEDICA – a Journal of Clinical Medicine 2016; 11(6): 122-129
- 20) Andreea-Iulia DOBRESCU, Adela CHIRITA-EMANDI, Nicoleta ANDREESCU, Simona FARCAS, Maria PUIU Does the Genetic Cause of Prader-Willi Syndrome Explain the Highly Variable Phenotype? MAEDICA – a Journal of Clinical Medicine 2016; 11(3):191-197
- 21) Dobrescu AI, Cosma M, Andreescu N, Farcaş S, Puiu M. Prader Willi Like syndrome- the new medical challenge Jurnalul Pediatrului Jurnalul Pediatrului, 2014, XVII (67-68):20-24.
- 22) Stoica F, Andreescu N, Olariu G, Jianu G, Puiu M. GENETIC POLYMORPHISMS AND RETINOPATHY OF PREMATURITY. Jurnalul Pediatrului, 2014, XVII (67-68):61-66.
- 23) RALUCA GRADINARU*, NICOLETA ANDREESCU*, MARIA PUIU CYP2D6 ENZYMATIC DEFICIENCY AND EXTRAPYRAMIDAL SIDES EFFECTS IN AN AUTISTIC PATIENT TREATED WITH RISPERIDONE Fiziologia - 2016.26.1(89):9-12
- 24) Anca Amalia Udriste, Natalia Cucu, Rodica Talmaci, Cosmin Arsene, Daniela Nedelcu, Maria Puiu, Alina Musetescu, Sabina Zurac, Nicoleta Andreescu, Octaviana Adriana Dulamea, Ileana Constantinescu, Marius NiculescuMethylation-specific PCR method for MGMT coding gene silencing evaluation and its prognostic significance in alkylating antitumor treatment Biointerface Research in Applied Chemistry, 2016, 6(6):1717-1721
- 25) Anca Amalia Udriste, Natalia Cucu, Vlad Constantinescu, Lilia Matei, Octaviana Adriana Dulamea, Ileana Constantinescu, Maria Mirela Iacob, Maria Puiu, Nicoleta Andreescu, Cosmin Arsene, Marius Niculescu. MS-MLPA method for the analysis of the glioma tumor MGMT encoding gene promoter methylation: treatment predictive considerations Biointerface Research in Applied Chemistry, 2016, 6(6):1737 – 1742
- 26) Simona Sorina Farcas, **Nicoleta Ioana Andreescu**, Andreea Iulia Dobrescu, Adela Emandi Chirita, Delia Hutanu, Florin Dorneanu, Natalia Usurelu, Victoria Sacara, Maria Puiu OBESITY IMPACT ON REPRODUCTIVE FITNESS Jurnalul Pediatrului, 2018, XXI (81-82):58-63
- 27) IE Jurca Simina, A Chirita-Emandi, **N Andreescu**, S Farcaş, A Mihailescu, AM Popa, P Tutac, C Zimbru, AI Dobrescu, IT Perva, A Murariu, M Puiu BURDEN OF RARE GENETIC DISEASES –EXPERIENCE OF TIMIS REGIONAL CENTRE OF MEDICAL GENETICS, ROMANIA Jurnalul Pediatrului, 2019, XXII (85-86):56-65
- 28) Iulia Maria Sabau, **Nicoleta Ioana Andreescu**, Adela Chiriță-Emandi, Iulia Jurca-Simina, Meda-Ada Bugi, Maria Puiu GENETICS IN ANOREXIA NERVOSA JURNALUL PEDIATRULUI – Year XXIV, Vol. XXIV, Nr. 93-94: 23-27
- 29) Iulia Maria Sabau, **Nicoleta Ioana Andreescu**, Adela Chiriță-Emandi, Iulia Jurca-Simina, Meda-Ada Bugi, Maria Puiu KETOGENIC DIET AND GENETIC DIS ORDERS JURNALUL PEDIATRULUI – Year XXIV, Vol. XXIV, Nr. 93-94: 28-33