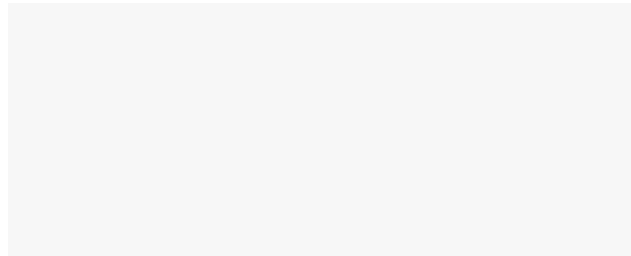


INFORMAȚII PERSONALE Puiu Maria



Din mai 2014 pana in prezent

Din mai 2010 pana in prezent

- Activitati de cercetare

Din octombrie 2009 pana in prezent

- Activitati de cercetare

Din octombrie 2003- oct. 2009

- Activitati de cercetare

Octombrie 1999 – octombrie 2003

- Activitati de cercetare

Martie 1992 – octombrie 1999

- Activitati de cercetare

EDUCAȚIE ȘI FORMARE

2010	Certificat de absolvire a modulului de formare in managementul universitar. Managementul cercetarii UE, POSDRU, AMPOSDRU, OIPOSDRU, Guvernul Romaniei, UEFISCDI Organizat in cadrul proiectului strategic Imbunatatirea Managementului Universitar, Cluj- Napoca, noiembrie 2010
	SC IDAS GROUP SRL <ul style="list-style-type: none">▪ Program de specializare/ Certificat de absolvire acordat de Ministerul Muncii si Ministerul Educatiei, Cercetarii si Tineretului
Iunie 2010	Manager proiect GRUPUL DE CONSULTANTA PENTRU DEZVOLTARE, BUCURESTI <ul style="list-style-type: none">▪ Program de specializare/ Certificat de absolvire acordat de Ministerul Muncii si Ministerul Educatiei, Cercetarii si Tineretului
2009	Manager proiect SC IDAS GROUP SRL <ul style="list-style-type: none">▪ Program de specializare/ Certificat de absolvire acordat de Ministerul Muncii si Ministerul Educatiei, Cercetarii si Tineretului
2005	Medic primar UMF „V. Babes” Timisoara, Spitalul clinic de urgență pentru copii „L. Turcanu” <ul style="list-style-type: none">▪ Genetica medicală
2002	Medic primar UMF „V. Babes” Timisoara, Spitalul clinic de urgență pentru copii „L. Turcanu” <ul style="list-style-type: none">▪ Pediatrie
2000	Medic specialist UMF „V. Babes” Timisoara, Spitalul clinic de urgență pentru copii „L. Turcanu” <ul style="list-style-type: none">▪ Genetica medicală
1995	Medic specialist UMF „V. Babes” Timisoara, Spitalul clinic de urgență pentru copii „L. Turcanu” <ul style="list-style-type: none">▪ Pediatrie
1994	Doctorand cu frecvență – Diploma Doctor Summa Cum Laude 03.02.1994 Prin Ordinul nr 6082 al Ministerului Învățământului Universitatea de Medicina și Farmacie “Carol Davila” București Titlul tezei: Patologia unor populații intens consangvinizate din Banat Coordonator: Prof. Dr. Constantin Maximilian
1991-1992	Medic de medicina generală Dispensar Comlos, Spital Jimbolia <ul style="list-style-type: none">▪ Medic pediatru
1988-1990	Medic de medicina generală Dispensar Brusturoasa, Spital Comanesti <ul style="list-style-type: none">▪ Medicina generală adulți
1985-1988	Medic stagiar Spitalul Municipal Timisoara <ul style="list-style-type: none">▪ Medicina generală▪

COMPETENȚE PERSONALE

Limba maternă Română

Alte limbi străine cunoscute

	INTELEGERE		VORBIRE		SCRIERE
	Ascultare	Citire	Participare la conversație	Discurs oral	
Limba franceza	B2	B2	B2	B2	B2
Certificat emis de Centrul Cultural Francez Timisoara					
Limba engleza	A2	A2	A2	A2	A2
Evaluare realizata in cadrul catedrei de limbi straine UMFT					

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

Competențe de comunicare

Capacitate de comunicare dobândita în cursul activitatii didactice si de cercetare, adepta a spiritului de echipa la locul de munca; conducator a peste 40 de lucrari de licenta.

Competențe organizationale/manageriale

Președinte executiv al Societății Romane de Genetica Medicală.

Coordonator Centru Regional de Genetica Medicala Timis

Colaborez de peste 10 ani cu Organizația Salvati copiii (din 2008 sunt membru în colegeul director), de peste 8 ani cu Asociația Prader Willi din Romania (director adjunct) și Asociația Williams Romania, sunt membru fondator și vicepreședinte al Aliantei Naționale a Bolilor Rare Romania (ANBRaRo). În aceasta calitate am organizat numeroase manifestări și campanii naționale pentru promovarea bolilor rare în Romania, implicand UMF Timisoara ca partener (6th International Prader-Willi Syndrome Scientific Conference and Rare Diseases Conference, Cluj, 2007, Conferința Națională cu participare internațională "Bolile rare – De la evaluarea nevoilor la stabilirea priorităților", Zalau, 2007, Seminarul : "Împreună pentru bolile rare", Timisoara, Rare day for rares diseases, februarie 2008, 2009, 2010, Simpozionul "Trust of Trust", Cluj, 2008, Conferința est europeana Prader Willi syndrome, 2009, Timisoara, Conferința balcanica de boli rare, 2009, Cluj).

Am participat la creionarea și implementarea Planului Național pentru Bolile Rare. Ca vicepreședinte ANBRaRo și specialist, particip la întâlnirile de lucru și la manifestări organize de Ministerul Sanatății Publice, Președinția României, Institutul Național de Sanatate Publică.

Am reprezentat România în foruri internaționale și sunt invitată la manifestări organize de acestea (Conferințe europene de Boli rare-2006, 2007, 2008), Meduse Conference (2007, Paris), EPOSSI Workshop (speaker, 2008), FRAMBU, Norvegia (2008) la manifestări internaționale.

Experiența în organizarea activitatii didactice și de cercetare la locul de munca (proiecte de cercetare în colaborare cu echipe multidisciplinare și multicentrice).

Am fondat și coordonez, în calitate de redactor șef Romanian Journal of Rare Diseases. În cadrul proiectului norvegiano-roman Noro, am organizat și coordonez activitățile E-Universității de Boli rare (<http://www.edubolirare.ro/index.html>)

Competențe dobândite la locul de munca

Consult și sfat genetic în sindroamele dismórfice, cromozomopatii. Stabilirea riscului de recurență în bolile genetice. Organizarea infrastructurii naționale pentru implementarea Planului Național pentru Bolile Rare. Organizarea și managementul Departamentului de Genetica al Spitalului clinic de urgență pentru copii „L. Turcanu” Timisoara. Am creat secția cu paturi pentru bolnavii cu afecțiuni genetice.

Competențe informatici

O bună stăpânire a instrumentelor Microsoft Office (absolvent curs Microsoft Project Advanced, 2010) Cunoștiințe ale aplicațiilor de grafică de calculator (Adobe Illustrator, PhotoShop)

Alte competențe	<p>Activitate de voluntariat si coordonator de voluntari (instruirea unui grup de voluntari, studenți la Facultatea de Medicina).</p> <p>Cultura organizanala si abilitate in scrierea si coordonarea proiectelor adaptate ONG cu activitate in Sanatate.</p> <p>Coordonez din 2007 un grup de studenți ai Universitatii de Medicina si Farmacie care desfasoara activitati complexe de voluntariat impreuna si pentru pacientii cu boli rare: Grupul "Voluntari pentru bolile rare". Impreuna cu acești studenți am scris si castigat numeroase proiecte iar activitatea studenților a fost apreciata in presa si in cadrul Galei Premiilor Carol Davila, unde a primit Premiu special.</p> <p>Premiu de Excelență acordat de Revista viata Medicală, 2010, pentru întreaga activitate în domeniul bolilor rare</p> <p>Premiu CMR, pentru MEDIC IMPLICAT, Gala Medica, Bucuresti, 2011</p>
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INFORMATII SUPLIMENTARE

Apartenenta la organizatii profesionale

1. Nationale:

- **2010-2018 Presedinte executiv Societatea Romana de Genetica Medicala**
- 2007 Alianța Națională pentru Boli Rare (membru fondator si vicepresedinte)
- Societatea Romana de Pediatrie
- Societatea Romana de Hematologie
- Societatea Romana de Pediatrie Sociala
- Societatea Romana de Biochimie si Biologie Moleculara,
- Asociatia Prader Willi din Romania (director adjunct)
- Organizatia Salvati copiii (membru in colegiul director)
- Asociatia Williams din Romania
- Coalitia Organizatiilor Pacientilor cu Afecțiuni Cronice (COPAC).

2. Internationale:

- American Society of Human Genetics (ASHG)
- European Society of Human Genetics (ESHG)
- European Cytogenetics Association (ECA)
- European Society for Clinical Investigation (ESCI)
- European Society for Medical Oncology (ESMO)
- Innovative Medicines Steering group (INNOMED-RO)

Publicații(1982-2012)

- Cursuri: 9, Indrumatoare: 8, Volume colective: 14, Monografii: 12
- Articole publicate in volume de rezumat la congrese internationale: 246
- Articole in extenso in reviste de circulatie nationala recunoscute: 234
- Articole publicate in volume de rezumat din tara: 261
- Lucrari comunicate in congrese si simpozioane nationale si internationale: 266.

Proiecte:

- Proiecte de cercetare: 16 (5 director, 1 manager, 1 asistant manager, 1 responsabil partener, 2 reprezentant specialisti, 4 coordonator specialist, 2 membru in echipa de cercetare).

ANEXE

LISTA LUCRARILOR REPREZENTATIVE

I. Monografii

1. **Maria Puiu** (coordonator), Medical Alert in Rare Genetic Diseases, Timisoara, "Victor Babes" Publisher, 2011, ISBN 606-8054-39-X;
2. **Maria Puiu** (coordonator) , Bolile rare, intre daruire si intelegerere, Ed. Brumar, Timisoara, 132 pag. ISBN 978-973-602-390-3, editura recunoscuta CNCSIS, 2008
3. **Maria Puiu** (coordonator) , Bolile rare, informatii utile pentru parinti, Ed. Brumar, Timisoara, 92 pag. ISBN 978-973-602-391-0, editura recunoscuta CNCSIS, 2008
4. **Maria Puiu** (coordonator), Esentialul in 101 boli genetice rare, Ed. Orizonturi Universitare, Timisoara, ISBN 978-973-638-327-4, editura recunoscuta CNCSIS, 512 pagini,2007
5. Cristina Rusu (coordonator), Metode uzuale in screeningul si diagnosticul bolilor genetice, Editura Gr. T. Popa U.M.F. Iasi, ISBN 978-973-7682-31-4, editura recunoscuta CNCSIS, 266 pg.; **Maria Puiu**: capitol 1.Tehnici de screening prenatal. Screeningul

- serului matern, pp 3-13, capitol 4. Tehnici de diagnostic prenatal. Amniocenteza, pp 71-81, Punctia de vilozitati corionice, pp 81- 86, Cariotipul, pp 105-112, 2007.
- 6. Mihai Gafencu, **Maria Julieta Puiu**, Violeta Stan, Gabriela Doroș, Sindromul Down de la îngrijire la înțelegere și acceptare, Ed. Brumar, ISBN 973-602-137-8, editura recunoscuta CNCSIS, 236 pagini, Maria Puiu, capitolul 1, pp 9-11, capitolul 2, pp 11-15, capitolul 3, pp 15-29, capitolul 12, pp 211-227, 2005.
 - 7. Mircea Covic, Dragoș Ștefănescu, Ionel Sandovici (coordonatori), Genetică medicală, Editura Polirom, Iași, ISBN 973-681-334-7, 607 pg, editura recunoscuta CNCSIS, xxx pagini, **Maria Puiu**, capitol 6. Variabilitatea genetica (M. Covic, I. Dimofte, M. Puiu, I. Sandovici), pp 203-248, capitol 7. Genetica populațiilor (M. Covic, M. Puiu, E. Severin), pp 253-271, 2004.
 - 8. **Maria Puiu**, Mic dicționar de genetică medicală, Ed. Eurobit, Timisoara, ISBN-973-9336-87-6, 210 pg, 1998
 - 9. **Maria Puiu**, Genetica populațiilor umane, Ed. Eurobit, Timisoara, ISBN 973-9336-86-8, 138 pg. 1998
 - 10. **Maria Puiu**, Genetica izolatorilor, Ed. Helicon, Timisoara, ISBN 973-9133-71-1, 173 pg. 1995

II. Capitole de carte

- 1. **Maria Puiu**, Adela Chirita Emandi and Smaranda Arghirescu (2013). Genetics and Obesity, Genetic Disorders, Maria Puiu (Ed.), p 271-292, ISBN: 978-953-51-0886-3, InTech, Available from: <http://www.intechopen.com/books/genetic-disorders/genetics-and-obesity>
- 2. **Maria Puiu**, Simona Dumitriu, Adela Chirita - mandi, Raluca Grădinaru and Smaranda Arghirescu (2013). The Genetics of Mental Retardation, Genetic Disorders, Maria Puiu (Ed.), p 143-174, ISBN: 978-953-51-0886-3, InTech, Available from: <http://www.intechopen.com/books/genetic-disorders/the-genetics-of-mental-retardation>
- 3. Ioana Micle si colab., Olimpia Tudose, **Maria Puiu**, Dorina Stoicanescu, *Diabetologie pediatrică – teorie si practica*, Capitolul, *Genetica diabetului zaharat tip 1*, Editura Marineasa, Timisoara, ISBN-973-9485-68-5, 2000.

III. Cursuri, indreptare lucrari practice

- 1. Maria Puiu, Dorina Stoicanescu, Gug Cristina, Simona Farcas, Popa Cristina, Nicoleta Andreescu, Adela Chirita-Emandi, Andreea Dobrescu; Aplicatii practice de Genetica pentru Asistenta Medicala generala, Editura "Victor Babes", Timisoara 2017, ISBN 987-606-786-044-3
- 2. **Maria Puiu**, D. Stoicanescu, C. Gug, S. Farcas, C. Popa, N. Andreescu, A. Chirita-Emandi, A. Dobrescu, Curs de Genetica Medicala, Ed. Eurostampa, Timisoara, ISBN 978-606-32-0296-4, 2016.
- 3. **Maria Puiu**, Genetica si farmacogenetica, Curs si lucrari practice pentru studentii facultatii de farmacie, Editura Brumar, Timisoara, ISBN 978-973-602-241-5, 209 pg. 2008
- 4. Valerica Belengeanu, **Maria Puiu**, Dorina Stoicanescu, Cristina Gug, Mirela Mihăescu, Simona Farcaș, Cristina Popa, Monica Stoian, Nicoleta Andreescu, Noemi Meszaros, Aplicatii practice in Genetica medicala, Ed. Eurostampa, Timisoara, ISBN 978-973-687-676-9, 272 pg, 2008
- 5. Valerica Belengeanu, **Maria Puiu**, D. Stoicanescu, C. Gug, M. Mihaescu, S. Farcas, C. Popa, M. Stoian, Elemente de Genetica medicala, Editura Orizonturi universitare, Timisoara, ISBN: (10) 973-638-272-9, 275 pagini, 2006
- 6. Valerica Belengeanu, **Maria Puiu**, Dorina Stoicanescu, Cristina Gug, Mirela Mihăescu, Simona Farcaș, Cristina Popa, Kinga Rozsnyai, Genetica medicala – Aplicații practice, Ed. Orizonturi universitare, Timișoara, ISBN 973-638-111-0, 160 pagini, 2004
- 7. **Maria Puiu**, Genetique medicale, cours et travaux pratiques, Editura Orizonturi Universitare, Timisoara, ISBN-973-8391-39-3, 2002
- 8. Olimpia Tudose, Valerica Belengeanu, **Maria Puiu**, Dorina Stoicanescu, Cristina Gug, Mirela Moga, Genetica medicala. CURS, Ed. Orizonturi universitare, Timișoara, ISBN 973-8109-09-4, 2000.
- 9. **Puiu, M.**, Moga, M, Notes de génétique médicale, Ed. Eurobit, Timisoara, ISBN 973-9441-88-4, 123 pg. Indexat NLM Catalog/PubMed, Notes de génétique médicale : à l'usage des étudiants en médecine de langue française, NLM ID: 101126830 [Book] , 1998.

LISTA PRINCIPALELOR LUCRARI PUBLICATE IN EXTENO

- 1. 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 doi: 10.1111/aos.13536
- 2. Ageu LŞ, Levai CM, Andreescu NI, Grigoraş ML, Hogea LM, Chiriac DV, Folescu R, Bredicean AC, Nussbaum LM, Enătescu VR, Poroch V, Lupu V, Puiu M, Nussbaum LA. 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

3. Adela Chirita Emandi, Diana Munteanu, Nicoleta Andreeescu*, Paul Tutac, Corina Paul, Iulian Puiu Velea, Agneta Maria Puszta, 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>
4. Dragoș Erdelean, Simona Sorina Farcăș, Vladimir Poroch, Nicoleta Ioana Andreeescu*, Izabella Erdelean, Andreea Iulia Dobrescu, Laura Alexandra Nussbaum, Lavinia Maria Hogea, 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
5. NCD Risk Factor Collaboration (NCD-RisC) Contributions of mean and shape of blood pressure distribution to worldwide trends and variations in raised blood pressure: a pooled analysis of 1018 population-based measurement studies with 88.6 million participants. . Int J Epidemiol. 2018 Mar 19.
6. Dumache R, Puiu M, Puszta AM, Parvanescu R, Enache A. A Single Step Mutation at D3S1358 Locus in a DNA Paternity Testing with 2 Alleged Fathers. Clin Lab. 2018 Sep 1;64(9):1561-1571.
7. Belei O, Olariu L, Puiu M, Jinca C, Dehelean C, Marcovici T, Marginean O Continuous esomeprazole infusion versus bolus administration and second look endoscopy for the prevention of rebleeding in children with a peptic ulcer. Rev Esp Enferm Dig. 2018 Jun;110(6):352-357
8. Serafim V, Shah A, Puiu M, Andreeescu 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.
9. Hogea LM, Nussbaum LA, Chiriac DV, Ageu LŞ, Andreeescu NI, Grigoraş ML, Folescu R, Bredicean AC, Puiu M, Roşca 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.
10. C Perva, IT Perva, DD Rusu, N Andreeescu, M Puiu Web based application for improving the education quality of young medical genetics healthcare professionals E-Health and Bioengineering Conference (EHB), 2017, 161-164
11. Cristian G Zimbru, Nicoleta Andreeescu, Adela Chirita-Emandi, Ioan Silea, Maria Puiu, Mihai D Niculescu Analysis of decision tree performance in predicting the relationship between a scored outcome and multiple single nucleotide polymorphisms E-Health and Bioengineering Conference (EHB), 2017, 57-60
12. CG Zimbru, N Andreeescu, A Chirita-Emandi, A Stanciu, Ioan Silea, Mihai D Niculescu, Maria Puiu Splice site pattern analysis and identification of similar sequences in the deep intron areas of human chromosome 21 E-Health and Bioengineering Conference (EHB), 2017, 145-148
13. NCD Risk Factor Collaboration (NCD-RisC). Worldwide trends in body-mass index, underweight, overweight, and obesity from 1975 to 2016: a pooled analysis of 2416 population-based measurement studies in 128·9 million children, adolescents, and adults. Lancet. 2017 Dec 16;390(10113):2627-2642.
14. NCD Risk Factor Collaboration (NCD-RisC). Worldwide trends in blood pressure from 1975 to 2015: a pooled analysis of 1479 population-based measurement studies with 19·1 million participants. Lancet. 2017 Jan 7;389(10064):37-55.
15. Chirita-Emandi A, Papa MC, Abrudan L, Dobrescu MA, Puiu M, Velea IP, Paul C. A novel method for measuring subcutaneous adipose tissue using ultrasound in children - interobserver consistency. Rom J Morphol Embryol. 2017;58(1):115-123
16. Aparaschivei, D., Todea, A., Păușescu, I., et al. (2016). Synthesis, characterization and enzymatic degradation of copolymers of ϵ -caprolactone and hydroxy-fatty acids. Pure and Applied Chemistry, 88(12), pp. 1191-1201.
17. NCD Risk Factor Collaboration (NCD-RisC). A century of trends in adult human height. Elife. 2016 Jul 26;5. pii: e13410.
18. RADU-IOAN URSU, NATALIA CUCU, GEORGETA-FLORENTINA URSU, ILEANA CRACIUNESCU, EMILIA SEVERIN, MARIA PUIU, LYGIA ALEXANDRESCU Frequency study of the FTO and ADRB3 genotypes in a Romanian cohort of obese children ROMANIAN BIOTECHNOLOGICAL LETTERS 2016, 21(3): 11610-11620
19. Chirita-Emandi A, Gabriela Barbu C, Cinteza EE, Chesaru BI, Gafencu M, Mocanu V, Pascanu IM, Tatar SA, Balgradean M, Dobre M, Fica SV, Ichim GE, Pop R, Puiu M: Overweight and Underweight Prevalence Trends in Children from Romania - Pooled Analysis of Cross-Sectional Studies between 2006 and 2015. Obes Facts. 2016 Jun 18;9(3):206–20.
20. Stoica F, Ladariu C, Koos MJ, Stanciu A, Olariu G, Andreeescu N, Puiu M. Refractive and Visual Outcome after Laser-Treated Retinopathy of Prematurity in Western Romania. Maedica (Buchar). 2016 Jun;11(2):122-129
21. Nicoleta Ioana Andreeescu, 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

22. Nicoleta Andreescu, Laura Nussbaum, Lavinia Maria Hoga, Raluca Grădinaru, Calin Muntean, Radu Ștefănescu, 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.
23. Laura Alexandra Nussbaum, Lavinia Maria Hoga, 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, 57(3):959–967
24. Laura Nussbaum, Nicoleta Andreescu, Lavinia Maria Hoga, Calin Muntean, Radu Stefanescu, Maria Puiu, Pharmacological and clinical aspects of efficacy, safety and tolerability of atypical antipsychotic medication in child and adolescents patients with schizophrenia and bipolar disorder, FARMACIA, 2016, 64(6):868-875
25. RALUCA CLAUDIA GRĂDINARU1, NICOLETA IOANA ANDREESCU, LAURA ALEXANDRA NUSSBAUM2, SIMONA SORINA FARCAŞ, VICTOR DUMITRĂSCU, LIANA SUCIU, MARIA PUIU-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):1343–1349
26. FLORINA STOICA, DANIELA IONESCU, ALINA HEGHES, CRISTINA TRANDAFIRESCU, NICOLETA ANDREESCU, ANCA TUDOR, SEBASTIAN OLARIU, ALINA STANCIU, MIHAELA GALEA, MARIA PUIU Vascular Endothelial Growth Factor Gene Polymorphism - Susceptibility Predictor for Severe Retinopathy of Prematurity? REV. CHIM.(Bucharest), 2016, 67(12): 2522-2525.
27. 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 Methylation-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
28. 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
29. 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
30. Chirita-Emandi A, Socolov D, Haivas C, Calapă A, Gheorghiu C, Puiu M. Vitamin D Status: A Different Story in the Very Young versus the Very Old Romanian Patients. PLoS ONE. 2015, 29;10(5):e0128010.
31. Adela Chirita-Emandi, Gabriela Doros, Iulia Jurca Simina, Mihai Gafencu, Maria Puiu, Head circumference references for school age children in western Romania, Rev. Med. Chir. Soc. Med. Nat., Iasi- 2015- Vol. 119, No. 4, page 1083-1091
32. RĂZVAN VLADIMIR SOCOLOV, NICOLETA IOANA ANDREESCU*, ANA MARIA HALICIU, EUSEBIU VLAD GORDUZA, FLORENTIN DUMITRACHE, RALUCA ANCA BALAN, MARIA PUIU, MIHAELA AMELIA DOBRESCU, DEMETRA GABRIELA SOCOLOV Intrapartum diagnostic of Roberts syndrome – case presentation Rom J Morphol Embryol 2015, 56(2):585–588 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
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PARTICIPAREA IN PROIECTE SI PROGRAME DE CERCETARE DEZVOLTARE

1. Proiect Science and Technology in childhood Obesity Policy (STOP), Grant Agreement number 774548; Call: Horizon 2020-SFS-2016-2017; Topic: SFS-39-2017, Research Innovation Action 2018-2020
2. Proiect cofinanțat din FONDUL SOCIAL EUROPEAN prin Programul Operațional Capital Uman 2014-2020 Axa priorităř: 4 Incluziunea socială și combaterea sărăciei - Obiectiv specific: 4.8 - Îmbunătățirea nivelului de competențe al profesioniștilor din sectorul medical; Titlul proiectului: Formarea PROfesională a personalului medical în GENetica medicală– PROGEN - SMIS 107623; Contract POCU: 91/4/8/107623/08.12.2017 (12.2017-12.2019) Coordonator implementare
3. Competitiveness Operational Programme 2014-2020; priority axis 1 – Research, technological development and innovation (RD&I) to support economic competitiveness and business development action 1.1.4 Attracting high-level personnel from abroad in order to enhance the RD capacity; Title: **Use of nutrigenomic models for the personalized treatment with medical foods in obese people (NutriGen)** 2016-2019 Coordonator implementare
4. Coordonator proiect: Centrul de Medicină Genomică v2; Universitatea de Medicină și Farmacie "Victor Babeș" Timișoara; ID: 1854; SMIS:487449 Nr.contract: 677/09.04.2015 POSCCE Operațiunea 2.2.1: Dezvoltarea infrastructurii CD existente și crearea de noi infrastructuri CD (laboratoare, centre de cercetare) Coordonator Proiect
5. Project „Quality Standards and Specific Performance Indicators for Health Education” POSDRU/18/1.2/G/40067POSDRU Expert calitate pe termen scurt
6. AXA PRIORITARĂ 1 “Educația și formarea profesională în sprijinul creșterii economice și dezvoltării societății bazate pe cunoaștere” DOMENIUL MAJOR DE INTERVENȚIE 1.5 “Programe doctorale și postdoctorale în sprijinul cercetării” Parteneriat interuniversitar pentru creșterea calității și interdisciplinarității cercetării doctorale medicale prin acordarea de burse doctorale – DocMed.net, 01.12.2010 - 30.11.2013 Expert pe termen lung
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| 6. Corelarea aspectelor clinice, genetice si epigenetice implicate in etiologia sindoamelor Prader Willi/Angelman: model de abordare multidisciplinara a bolilor rare in Romania, PNCD, Program Parteneriate, contract 42113, 2008-2011 | Director de proiect |
| 7. Monitorizarea bolii minime reziduale in leucemii acute limfoblastice la copil prin citometria in flux Multiparametrica, CNCSIS tip A, 2007-2008 | Manager proiect |
| 8. Optimizarea managementului copiilor cu LAL prin folosirea tehnicilor de citogenetica moleculara (FISH) in protocolul de evaluare, CNCSIS tip A, 2007-2008 | Director de proiect |
| 9. Optimizarea diagnosticului si managementului pacientilor cu retard mintal prin introducerea in protocolul de evaluare a testului MLPA, CNCSIS cod 832, 2006-2007 | Responsabil partener |
| 10. Romanian National Alliance for Rare Diseases – RONARD, Trust for Civil Society for CEE, RO/IX 2006/123, 2007-2008 | Coordonator specialisti,
Director adjunct |
| 11. Corelatii intre distributia parenchimatoasa a elementelor de angio-si bilioarhitectura si segmentarea lobului caudat - Baze anatomice pentru chirurgia de rezectie si transplant, PNCDI 2 – Program 4 – Parteneriate, cod 2167/2007-2009 | Membru in echipa de cercetare |
| 12. Împreună pentru o viață mai bună pentru pacienții cu boli rare din România, Matra Kap Programme, finantare Ambasada Regatului Țărilor de Jos, 2005-2006 | Coordonator specialisti |
| 13. Rare Diseases Solidarity Project, Romanian National Alliance for Rare Diseases (RONARD), The Trust for Civil Society in Central & Eastern Europe ("CEE Trust") RO_X 2007_190, October 2008 - September 2009 | Assistant manager |
| 14. Programul National al MSP: PII/9: Managementul Registrelor de boli cronice la copil, 2008. | Coordonator proiect |
| 15. Programul National al MSP: PII/9: Diagnosticul genetic al Miodistrofiilor Duchenne si Becker, 2009. | Coordonator proiect |
| 16. Proiect NoRo - finantat de Innovation Norway, parteneri APWR, UMFT, Ministerul Sanatatii Publice, 2008-2011 | Director adjunct,
Responsabil partener
UMFT |
| 17. TREAT-NMD Neuromuscular Network “Accelerating Treatments for Neuromuscular Diseases” | Membru |