

CURRICULUM VITAE



GENERAL INFORMATION

Name

Martino (Michele Lucio Giovanni) RUGGERI

Address

DEPARTMENT OF CLINICAL AND EXPERIMENTAL MEDICINE
SECTION OF PEDIATRICS AND CHILD NEUROPSYCHIATRY
UNIVERSITY OF CATANIA

[HTTPS://WWW.MEDCLIN.UNICT.IT/DOCENTI/MARTINO.RUGGERI](https://www.medclin.unict.it/docenti/martino.ruggieri)

AZIENDA OSPEDALIERO-UNIVERSITARIA "POLICLINICO" (AOUP)

VIA S. SOFIA, 78 - CATANIA - PO "G. RODOLICO"

BUILDING 3, FLOOR 1

UNIT OF PEDIATRIC CLINIC

POSTGRADUATE TRAINING PROGRAM IN PEDIATRICS

REGIONAL REFERRAL CENTRE FOR "RARE DISEASES OF THE NERVOUS SYSTEM IN CHILDHOOD" AND "EXPANDED NEWBORN SCREENING AND INHERITED METABOLIC DISEASES"

METABERN [METABOLIC EUROPEAN RARE NETWORK] – CATANIA, ITALY

Phone number

++39 095 3781193/3782394 [Unit of Pediatric Clinic, Ward]

Mobile: ++39 338 5084769

Fax

++39 095 3782940

E-mail

m.ruggieri@unict.it; clinicapediatricacatania@gmail.com;
malattierarecatania@gmail.com

Nationality

Italian

Birth date / Fiscal code

21ST APRIL 1962 [CATANIA] / RGGMTN62D21C351E

POSITIONS HELD

- From [from - to]
- Name and Address
- Department
- Type of job

* From 29th December 2015 - up to now

University of Catania

Department of Clinical and Experimental Medicine

Professor and Chairman of Pediatrics (MEDS/20A)

[Rectorial Decree no. 164393 dated 29/12/2015]

• **Positions held**

* From 8th October 2019 - up to now

Deputy Rector "*International Affairs*" [Biomedical area]
[Rectorial Decree no. 2984 dated 08/10/2019]
University of Catania

* From 1st November 2022 - up to now

Director - Unit of Pediatric Clinic
AOU "*Policlinico*", PO "*G. Rodolico*"
[Resolution of the General Director no. 2360 dated 2/11/2022]
University of Catania

* From 17th October 2018 - to 31st October 2022

From 1st November 2022 - up to now [2nd term]

Director, Postgraduate Training Program in Pediatrics
[Rectorial Decree no. 137929 17/10/2018]
University of Catania

* From 7th January 2022 - to 31st December 2024

President

Italian Society of Pediatric Neurology (Società Italiana di Neurologia Pediatrica, **SINP**)
Currently - **Past-President**

* From 1st December 2023 - up to now

Delegate - Committee of the National Scientific Societies (SIP)
Member - Board of the Italian Society of Pediatrics (SIP)

* From 19th September 2016 - to 30th October 2022

Head - Unit of Rare Diseases of the Nervous System in Childhood
AOU "*Policlinico*", PO "*San Marco*"
[Resolution of the General Director no. 1555 dated 19/09/2016]
University of Catania

* From 10th November 2019 - to 25th May 2022

Member - Coordination Committee "*Faculty of Medicine and Surgery*"
[Representative of the Postgraduate Training Programs in Medicine]
University of Catania

* From 1st May 2014 - to 8th February 2020

Member, Local Ethics Committee - Catania 1 - Catania
AOU "*Policlinico*" - Catania
University of Catania

* From 30th October 2016 - to 14th October 2018

Vicedirector - Postgraduate Training Program in Pediatrics
University of Catania

* From 23rd February 2012 - to 31st October 2014

Vicedirector - Department of Educational Sciences [**DISFOR**]
[Rectorial Decree no. 159/18 dated 23rd February 2012]
University of Catania

PREVIOUS POSITIONS [Italy & Abroad]

* From 15th January 2015 - to 29th December 2015

University of Catania

Department of Clinical and Experimental Medicine

Associate Professor of Pediatrics (MED/38)

* From 15th November 2009 - to 14th January 2015

University of Catania

Department of Educational Sciences [DISFOR]

Associate Professor of Pediatrics (MED/38)

* From 23rd February 2012 - 31 ottobre 2014

Vicedirector – Department of Educational Sciences [DISFOR]

[Rectorial Decree DISFOR no. 159/I8 23/02/2012]

University of Catania

* From 1st November 2006 - to 14th November 2009

National Research Council (CNR)

Institute of Bioimaging and Pathophysiology of the Nervous System (IBFSNC)

→ Institute of Neurological Sciences / Department of Pediatrics

University of Catania

1st Researcher in Pediatrics (MED/38)

* From 23rd August 2000 - to 31st October 2006

National Research Council (CNR)

Institute of Bioimaging and Pathophysiology of the Nervous System (IBFSNC)

→ Institute of Neurological Sciences / Department of Pediatrics

University of Catania

Researcher in Pediatrics (MED/38)

* From 1st May 1995 - to 31st March 1999

University of Oxford/National Health System (NHS)

Oxford University Hospitals (OUH)

John Radcliffe Hospital/Churchill Hospital/Radcliffe Infirmary

Department of Pediatrics (Pediatric Neurology), Clinical Genetics and Neuroradiology

Clinical Assistant (and Research Fellow) with Senior Registrar Status

→ Recognised by the Italian Ministry of Health - Decree no. 0012008-P-DGPROF_02-03-2020 del 2 Marzo 2020

- From 1st November 1995 to 31st May 1998;

- From 6th July to 10th September 1998

- From 15th to 26th March 1999

* From 1st July 1995 - to 1st November 1995

University of Harvard, Boston, USA

Massachusetts General Hospital (MGH)

Department of Pediatrics, Neurology and Neurogenetics

Clinical Research Fellow

* From 22nd April 1992 - up to now

[Certified Italian General Medical Council no. 10300]

EDUCATION / TRAINING

• From [from – to]	* From 1st November 1995 - to 1st March 2000
Name and type of Institution	University of Catania/University of Oxford
• Type of courses	Pediatric Sciences
• qualification awarded/achieved	PhD [XI cycle] in Pediatric Sciences [achieved on 26th February 2000]
	* From 1st April 1992 - 31st October 1995
	University of Catania
	Department of Pediatrics
	Resident in Pediatrics
	Residency in Pediatrics achieved on 9th November 1995 [50/50 cum laude]
	* From 1st November 1982 - to 31st October 1984
	University of Oxford, GB
	Baccalaureate in <i>"History of Modern and Contemporary British Literature"</i>
	BA history [Thesis: Life and novels of Graham Greene]
	* From 1st November 1982 - to 5th July 1991
	University of Catania
	Degree in Medicine and Surgery (110/110 cum laude)
	M.D. in Medicine and Surgery
	* From 1st October 1976 - to 31st July 1981
	Institute "Salesiani" di San Francesco di Sales" - Classical Studies"
	Catania
	Secondary School II Level - Classical Studies
	High School Diploma (60/60 cum laude)
	* From 1st October 1973 - 9th June 1976
	Institute "Salesiani" di San Francesco di Sales" – Classical Studies"
	Catania
	Secondary School I Level - Classical Studies
	* From 1st October 1968 - 10th June 1973
	Institute "S. Giuseppe", Suore Maria Ausiliatrice
	Institute Salesiani "San Francesco di Sales"
	Primary School

NATIVE LANGUAGE	ITALIAN
FURTHER LANGUAGES	ENGLISH [excellent] [excellent] [excellent] Proficiency C2 level; School, Graduate and post-graduate periods in English speaking Countries [1976, Norwich; 1977, Oxford; 1978, Oxford; 1979, Oxford; 1982-1984, Oxford (BA History of Modern and Contemporary British Literature); 1995-2000, Oxford [John Radcliffe Hospital, University of Oxford]; 1996, Boston (Massachusetts General Hospital, Harvard Medical School)] FRENCH [excellent] [excellent] [excellent] DELF B2 level [1974, Nizza; 1975, Monaco] SPANISH [excellent] [excellent] [excellent] DELE C1 level [1980; Barcellona/Eurocenter; 1981, Vigo/Eurocenter] GERMAN [basic] [basic] [basic]
FURTHER COMPETENCIES.	General Medical Council Certificate [(GMC) London, GB, June 1995, no. 4179737]
LICENCES	DRIVING LICENCE B NAUTICAL LICENCE [From December 2006]

SCIENTIFIC AWARDS / PRIZES

- Prize "Felice Paradiso" for Pediatric Theses in Pediatric Science (June 1994)
- Prize "Kiwanis" for excellency in Medicine and Surgery (March 1995)
- V Prize "Paolo Balestrazzi" for the Study of the neurofibromatoses, Venafrò (May 2002)
- Prize Militello - Firenze degli Iblei – for his pediatric studies, Militello Val di Catania (CT) (Ottobre 2010)
- "Robert J. Gorlin" Award Prize [Robert Gorlin Memorial Lecture] - AAOMP [American Academy of Oral and Maxillo-Facial Pathology] - Chicago, USA (Virtual conference) (24 May 2021)

https://mediaspace.umn.edu/media/t/1_ftthonxlt

GRANTS

CNR, A196.00174.04 (Roma, 1996): "Studio clinico e genetico della NF1 segmentale" (5 million of Liras)
Oxfordshire Health Services Research (Oxford, GB 1996-98): "Segmental NF1" (10,000 GBP)
Medical Research Council (Oxford, 1997-1998): "Spinal lesions in the neonatal age" (5,000 GBP)
"ASSERT" Angelman Lay Group (Londra, 1996-1998): "Epilepsy in the setting of Angelman syndrome" (5,000 GBP)
Italian Ministry of Health [Ministero della Salute], Regional Projects 2004-2008: "Oncogenes in NF1" (210,000 euros)
National Institute of Health (Boston, 2003-2008): "Pediatric Multiple Sclerosis" (510,000 USD)
CNR, Commessa Ricerca a Tema Libero (Rome, 2006): "Sindromi neurocutanee" (12,000 euro)
CNR, Commessa Ricerca a Tema Libero (Rome, 2008): "Le neurofibromatosi" (13,000 euro)
Tuberous Sclerosis Fundings AST onlus (Rome, 2012-2013) "The role of mi-RNA in tuberous sclerosis" (12.000 euros)
Tuberous Sclerosis Fundings AST onlus (Roma, 2014-2015) "The role of mi-RNA in tuberous sclerosis" (15.000 euros)
FIR 2014-2015, University of Catania. Project 922B27 - "Puer Sapiens" Società, cultura e educazione infantile nella preistoria protostoria in Sicilia: analisi e realizzazione di modelli in 3D in scala reale" (12.000 euros)
ANAT (Associazione Nazionale Atassia-Telangiectasia) 2021-2022 Prize Best Project 2021 (15.000 euros)
Interdipartimental Project of the University of Catania 2020-2022 - Ricostruzione 3D di encefali [TC/sincrotrone] e Paleo-fenotipi e studio genomico [Y, NGS, mitocondrio, WES] in 3 resti fossili infantili Paleolitici [13.000 a.C. "Grotta dell'Uzzo" - Museo Salinas, PA] (45.000 euro)
POS (Progetti Operativi Sanitari T4 (Traiettorie T4 (February 2022 - December 2027), "Riposizionamento di farmaci nelle malattie rare neurologiche in età pediatrica e creazione di Hun e Spoke nelle malattie Rare" [coordinatore unico di Ateneo] (Unict, 12.500.000 euros)

EDITOR-IN-CHIEF / ASSOCIATE EDITOR (Mamber EDITORIAL BOARD

2023 - up to now Editor-in-Chief: Perspectives in Pediatric Neurology [PPN], Edizioni Mattioli1885

www.mattioli1885.com

2023 – up to now Associate editor: Frontiers in Human Neurosciences

www.springernature.com

2005 – 2011 Deputy Editor: Journal of Brachial Plexus and Peripheral Nerve Injury

www.jbpni.com

2007 - 2011 Associate Editor: Child's Nervous System

www.springer.com

2014 - up to now Associate Editor: Multiple Sclerosis and Other Demyelinating Disorders

www.msddjournal.com

2012 - up to now Advisory Editor: The Child

www.thechild.it

2012 - up to now Comitato Scientifico: Prospettive in Pediatria [Rivista Ufficiale della SIP]

www.prospettiveinpediatria.it

2014 - up to now Editorial Board: Behavioural Neurology

PRESIDENT of SCIENTIFIC SOCIETIES

President: Società Italiana di Neurologia Pediatrica (SINP) [7 January 2022 - 31 December 2024]

Secretary: Gruppo di Studio di Neuroimmunologia Pediatrica (GNIP) www.sinp.it

President: Comitato Scientifico, Associazione Neurofibromatosi (ANF), Parma [2017 - 2023]

PhD courses

Member of Board of Teachers: Dottorato di Ricerca in "Biomedicina Traslazionale" XXVIII ciclo, XXIX ciclo, XXX ciclo, XXXI - Università degli Studi di Catania (dal 2012 - ad oggi)

Member of BOARDS - SCIENTIFIC COMMITTEES

1998 - Comitato Scientifico, Associazione Nazionale Neurofibromatosi, A.N.F.
<http://www.neurofibromatosi.org>
<http://www.sclerosituberosa.it>
2000 - 2010 Italian Delegate "Committee of National Advisers" (CNA) in Paediatric Neurology, European Pediatric Neurology Society (EPNS)
<http://www.epns.com>
2002 - 2008 Società Italiana Neurologia Pediatrica, S.I.N.P.
2002 - International Scientific Committee "Hypomelanosis of Ito/HITS, UK <http://www.e-fervour.com/hits/#support>
2004 - Comitato Scientifico, Associazione Neurofibromatosi "Io ci sono" (BO)
<http://www.associazioneiocisono.com>
2004 - Scientific Consultant "Sclerosi Multipla infantile" Serono S.R.L.
www.serono.com
2009 - International Pediatric Multiple Sclerosis Study Group (IPMSSG)
<http://www.ipmssg.org>
2014 - 2018 Consiglio Direttivo nazionale, Gruppo di Studio di Storia della Pediatria (GSSP/SIP)
www.sip.it
2015 - 2018 Consiglio Direttivo, Società Italiana di Ricerca Pediatrica (SIRP)
www.sirp.it
2023 – ad oggi Consiglio Direttivo, Società Italiana di Pediatria (SIP)

EDITOR INTERNATIONAL BOOKS or TREATISES

Ruggieri M, Pascual-Castroviejo I, Di Rocco C.
Neurocutaneous diseases: Phacomatoses and hamartoneoplastic syndromes
New York-Wien: Springer-Verlag, 2008 (72 capitoli, 1040 pagine, 457 figure colori/B&N)
<http://www.springer.at>
ISBN: 9783211213964

EDITOR NATIONAL BOOKS / AUTHOR CHAPTERS

Ruggieri M
Neurologia Pediatrica – dalle basi biologiche alla pratica clinica
Milano: EDRA/Elsevier, 2023 (830 pagine)
<http://www.edra.com>
ISBN: 978-88-214-4865-2

Pavone L, Ruggieri M.
Neurologia Pediatrica.
Milano: Elsevier/Masson, 2^a edizione, 2006, 850 pagine
<http://www.masson.it>
ISBN: 88-214-2789-7/978882142789-3

Pavone L, Ruggieri M.
Neurologia Pediatrica.
Milano: Elsevier/Masson, 2001 (608 pagine)
<http://www.masson.it>
ISBN: 88214-2618-1

Ruggieri M, Tenconi R.
Le neurofibromatosi.
Edizioni A.N.F., Parma, 2000 (90 pagine) (2° edizione, 2007, 200 pagine)
<http://www.neurofibromatosi.org>

Ruggieri M, Migone N.
Sclerosi Tuberosa
Edizioni A.S.T., Roma 2008 (80 pagine)
<http://www.sclerosituberosa.org>

Ruggieri M, Franzoni E.
Neurologia e Psichiatria dello Sviluppo.
Milano: Elsevier/Masson (360 pagine) (2012)
<http://www.elsevier.com>
ISBN: 97888821-432682

Catassi C, Cogo P, Corsello G, Iughetti L, Peroni D, Piacentini G, Ruggieri M, Verrotti A.
Lissauer T, Carroll W. Manuale di Pediatria
Milano: EDRA, 2018: pp. 123-144
ISBN: 788821-4447877

Reviews online
Ruggieri M, Pavone L.
Hypomelanosis of Ito & related disorders
San Diego MedLink Neurology Database
www.Medlink.com

PRINCIPAL INVESTIGATOR Scientific Projects

Clinical, genetic and neuroradiological study of neurocutaneous syndromes
CNR project, 2000-2004
Study of oncogenes in neurofibromatosis type 1
MURST-Molise Region project, 2004-2008
SINP Group Database Childhood Multiple Sclerosis
Collaboration of the Italian Society of Neurology (SIN)
I study immunological and cellular factors Infantile Multiple Sclerosis
Hospital for Sick Children, Toronto, Canada/NIH 2003-200
POS (Health Operational Projects T4 (Trajectory T4
(October 2022-December 2025), "Repositioning drugs in neurological rare diseases at age
pediatric and creation of Hun and Spoke in Rare" diseases [unique coordinator of the University]
(Unict, EUR 12 500 000)

Studies conducted with GCP [Good Clinical Practice]

- Medical Research Council (Oxford, 1997-1998): "Lesioni midollo spinale neonatali"
RISULTATI pubblicati su: Dev Med Child Neurol 1999;41:51-54
- "ASSERT" Angelman Lay Group" (Londra, 1996-1998): "Epilessia ed Angelman"
RISULTATI pubblicati su: Arch Dis Child 1998;79:423-426
- Ministero della Salute, Progetti regionali 2004-2008: "Oncogeni nella NF1"
RISULTATI pubblicati su: Hum Mutat 2004;23:134-146; Hum Mutat 2008;29:74-82
- National Institute of Health (Boston, 2003-2008): "Sclerosi multipla infantile"
RISULTATI pubblicati su: The Lancet Neurology 2007;6:773-781; 112;
J Neuroimmunol 2010;223:92-99

REFEREE in peer-reviewed journals

The New England Journal of Medicine (Boston, USA)
The Lancet (London, UK)
The Lancet Neurology (London, UK)
Neurology (Rochester, USA)
Genetic Epidemiology (Rochester, USA)
Journal of Medical Genetics (Birmingham, UK)
Pediatrics (St Paul, USA)
Journal of Pediatrics (Cincinnati, USA)
Journal American Academy Dermatology (Chicago, USA)
Journal of Investigative Dermatology (Boston, USA)
American Journal Medical Genetics (Salt Lake City, USA)
Archives of Dermatology (Chicago, USA)
Developmental Medicine Child Neurology (London, UK)
Journal of Neurological Science (Chicago, USA)
European Journal of Neurology (Paris, FRANCE)
Archives of Disease in Childhood (London, UK)
European Journal Pediatric Neurology (Leuven, B)
Dermatology (Nice, F)
Pediatric Neurology (Rochester, USA)
Neuropediatrics (Essen, D)
Clinical Anatomy (Rochester, USA)
Acta Paediatrica (Stockholm, SW)
European Journal of Pediatrics (Zurich, CH)
Child's Nervous System (Rome, IT)
Case Reports and Clinical Practice Review (Warsaw, PL)
Journal Clinical Experimental Medicine (Warsaw, PL)
Journal of Pediatric Neurology (Van, TURKEY)
Neural Regeneration Research (NRR) (Shanyeng, CHINA)

SPECIAL MENTION

Award "Robert J. Gorlin" - Memorial Lecture, American Academy of Oral and Maxillofacial Pathology [AAOMP], New Orleans, USA - April 25, 2020 Reading "Neurocutaneous Disorders: From the People Behind Syndromes to Molecular Pathways and Targeted Therapies" [Salt Lake City, USA - May 24, 2021].

A complex malformation syndrome with mental retardation, dysmorphic signs and skin anomalies (cutis tricolor) is called Ruggieri-Happle syndrome

[Happle R. Mosaicism of human skin. Berlin: Springer-Verlag, 2014: Ruggieri-Happle syndrome; Torchia et al. Cutis 2013;91:11-16; Tekin B, et al. Dermatol Online 2014; 20(10)]

POSSUM syndrome 6275 MCA, Ruggieri-Happle syndrome (<http://www.possum.net.au>)

Published in: Eur J Pediatr 2000;159:745-749 and in Am J Med Genet 2003;120A:110-116)

A complex malformation syndrome with mixed vascular cutaneous nevus and brain anomalies of the Dyke-Davidoff-Masson type bears the name Ruggieri-Leech syndrome

[Happle R. Mosaicism of human skin. Berlin: Springer-Verlag, 2014: Ruggieri-Leech syndrome]

Published in: Am J Med Genet 2012;150A:1870-1880

A malformative syndrome with cardiac venous return anomaly (vein "scimitar" anomaly), multiple cardiac malformations and dysmorphic notes of the skull and face is called scimitar (vein) syndrome, such as Ruggieri (Vena Ruggieri Scimitar)

POSSUM 6256 MCA syndrome, Ruggieri type (scimitar vein abnormality, multiple cardiac malformations, craniofacial abnormalities) (<http://www.possum.net.au>)

LDDB and LNDB. Baraitser M & R Winter. Oxford: Oxford University Press, 2005

Published in: Am J Med Genet 2003;116A:170-175

INTERNATIONAL RESEARCH PAPERS [Scopus, WOS, PubMed]

1994 - 2025 (August)

IMPACT FACTOR (IF) ** TOTAL = 1.640

[** n. 383 peer-reviewed appearing on Scopus/WOS/PubMed con IF]

IF = from the Journal Citation Reports, JCR (ISI), 2025 - <https://www.jcrweb.com>

IMPACT FACTOR (IF) AVERAGE = 3.545

Total number of research papers Scopus/WOS = 426

H-index [Scopus; WOS; Researchgate] = 58

Total number of citations [Scopus; Researchgate] = 9.800

INTERNATIONAL RESEARCH ARTICLES [summary]

[Scopus - WOS - PubMed] → 1994 - 2025 (August)

1. Int J Ped Otorhinolar	1994;30:79-84	IF = 1.583	
2. Dev Brain Research Dysf	1994;17:20-28.	IF = 0.125	
3. Neuropediatrics	1994;25:1	IF = 1.947	
4. Ital J Pediatr	1994;20:549-553	IF = 2.638	
5. Neuroradiologia	1994;1:377-381		[6.290]
6. Pediatr Radiol	1995;25:34-36.	IF = 2.505	
7 Am J Med Genet	1995;59:139-142.	IF = 2.802	
8. Pediatr Radiol	1995;25:S1:147-149.	IF = 2.505	
9. Ital J Pediatr	1995;21:88-96	IF = 2.683	
10. Ital J Pediatr	1995;21:743-746	IF = 2.683	[13.173]
11. Am J Med Genet	1996;61:178-181.	IF = 2.802	
12. Genes Chromos Cancer	1996;15:18-25.	IF = 5.006	
13. Eur Heart J	1996;17:968	IF = 22.673	
14. Neurology	1996;45:485-492.	IF = 9.901	
15. Clin Dysmorph	1996;5:223-229.	IF = 0.813	
16. Clin Pediatr	1996;35:209-212.	IF = 0.840	
17. Clin Pediatr	1996;35:365-365.	IF = 0.840	
18. J Neurosurg	1996;85:941-944.	IF = 2.279	[44.916]
19. Curr Pediatr	1997;7:167-176	IF = 0.345	
20. Ped Dermatol	1997;14:22-25.	IF = 1.588	
21. J Med Genet	1997;34:256-260.	IF = 6.318	
22. J Neuroimmunol	1997;76:189-192.	IF = 3.458	
23. Am J Med Genet	1997;71:271-274.	IF = 2.802	
24. Clin Pediatr	1997;36:529-534.	IF = 0.840	
25. Clin Dysmorph	1997;6:375-378.	IF = 0.816	
26. Ital J Pediatr	1997;23:111-117	IF = 2.683	
27. Genet Counsel	1997;8:353-354	IF = 2.537	
28. Genet Counsel	1997;8:367-368	IF = 2.537	
29. Genet Counsel	1997;8:374-375	IF = 2.537	[26.461]
30. Pediatrics	1998;101:112-119.	IF = 5.800	
31. Br J Radiol	1998;71:225-228.	IF = 1.814	
32. Hum Genet	1998;102:591-597.	IF = 2.690	
33. Postgr Med J	1998;74: 257-259.	IF = 2.078	
35. Arch Dis Child	1998;79:423-426	IF = 3.258	
35. Ital J Pediatr	1998;24:XVII-XVIII	IF = 2.683	[23.133]
36. Dev Med Child Neurol	1999;41:51-54.	IF = 5.449	
37. Or Surg Or Med Or Pathol	1999;87:67-72.	IF = 2.589	
38. J Pediatr Orthop	1999;19:301-305.	IF = 1.010	
39. Dev Med Child Neurol	1999;41:311-317.	IF = 6.449	
40. Acta Paed	1999;88:671-674.	IF = 1.634	
41. Neurology	1999;88:671-674.	IF = 9.901	
42. Minerva Pediatr	1999;51:395-8	IF = 1.312	
43. Neurol Sciences	1999;20:89-108.	IF = 3.181	
44. Child's Nerv Syst	1999;15:295-308.	IF = 1.465	[32.000]

45. J Med Genet 2000;37:44-49	IF = 6.138	
46. Arch Dermatol 2000;136:426-427	IF = 2.339	
47. Eur J Pediatr 2000;159:477-480	IF = 2.335	
48. J Neurosurg 2000;93:530-532	IF = 2.279	
49. Eur J Pediatr 2000;159:745-749	IF = 2.335	
50. Am J Med Genet 2000;95:82-84	IF = 2.802	
51. Pathol Res Pract 2000;196:713-718	IF = 3.250	
52. Virchow Arch 2000;437:401-412	IF = 2.848	
53. Minerva Pediatr 2000;52:357-366	IF = 1.312	
54. J Child Neurol 2000;15:635-644	IF = 1.987	[27.270]
55. Neurology 2001;56:827-829	IF = 9.901	
56. Neurology 2001;56:1433-1443	IF = 9.901	
57. The Lancet 2001;357:311-312	IF = 79.321	
58. Arch Pathol Lab Med 2001;125:599-601	IF = 5.534	
59. Pediatr Neurol 2001;24:300-302	IF = 3.372	
60. Am J Med Genet 2001;101:178-180	IF = 2.802	
61. Neurology 2001;56:1606-1607	IF = 9.901	
62. Clin Neurol Neurosurg 2001;103:151-154	IF = 1.876	
63. Eur J Pediatr Neurol 2001;5:167-168	IF = 3.140	[125.378]
64. Virch Arch 2002;441:525-526	IF = 2.848	
65. Arch Gerontol Geriatr 2002;suppl 8:157-163	IF = 3.250	
66. Arch Gerontol Geriatr 2002;suppl 8:295-301	IF = 3.250	
67. Arch Gerontol Geriatr 2002;suppl 8:303-308	IF = 3.250	
68. Arch Gerontol Geriatr 2002;suppl 8:309-312	IF = 3.250	
69. Arch Gerontol Geriatr 2002;suppl 8:313-317	IF = 3.250	
70. Arch Gerontol Geriatr 2002;suppl 8:319-326	IF = 3.250	[22.348]
71. Am J Med Genet A 2003;116A:170-175	IF = 2.802	
72. J Med Genet 2003;40:227-232	IF = 6.138	
73. Am J Med Genet A 2003;120A:110-116	IF = 2.802	
74. Case Rep Clin Pract Rev 2003;4:2-5		
75. Ital J Pediatr 2003;29:222-225	IF = 2.683	[14.380]
76. Neurol Sciences 2004;25(suppl4):S326-335	IF = 3.181	
77. Hum Mutat 2004;23:134-46	IF = 4.868	
78. Br J Ophthalmol 2004;88:1429-1433	IF = 4.638	
79. Neuropediatrics 2004;35:207-210	IF = 1.205	
80. Neurol Sci 2004;25(suppl4):S346-349	IF = 3.181	[12.560]
81. Neuropediatrics 2005;36:21-34	IF = 1.205	
82. Am J Med Genet 2005;136A:357	IF = 2.802	
83. Acta Paediatr 2005;94:1066-1072	IF = 1.634	
84. Neuropediatrics 2005;36:279-283	IF = 1.205	
85. J Pediatr Endocrinol Metab 2005;18:1019-1025	IF = 1.634	
86. Ital J Pediatr 2005;31:280-283	IF = 2.683	[10.273]
87. Pediatr Neurol 2006;34:66-71	IF = 3.372	[3.372]
88. The Lancet Neurol 2007;6:773-81	IF = 44.182	
89. Pediatric Neurology 2007;36:607-610	IF = 3.372	
90. Am J Gastroenterol 2007;102:1831	IF = 10.843	
91. Pediatr Neurol 2007 Sep;37(3):209-11	IF = 3.372	
92. Am J Hum Genet 2007;81:104-113	IF = 11.025	[72.815]

93. J Pediatr 2008;152:244-249	IF = 4.806	
94. J Pediatr 2008;153:298-299	IF = 4.806	
95. Hum Mutat 2008;29:74-82	IF = 4.878	
96. Epilepsy Res 2008;78:225-31	IF = 2.740	
97. J Clin Gastroenterol 2008;42:715-9	IF = 3.062	
98. Minerva Pediatr 2008;60:383-4	IF = 1.312	
99. Clin Genet 2008;74:164-70.	IF = 4.378	
100. J Pediatr Hematol Oncol 2008;30:628-30	IF = 1.060	
101. Minerva Pediatr 2008;60:1473-4.	IF = 1.312	
102. Neurol Sci 2008;29:495-6	IF = 3.181	
103. Orthopaedics 2008;31:498	IF = 1.390	[31.291]
104. Br J Ophthalmol 2009;93:175-176	IF = 3.384	
105. Childs Nerv Syst 2009; 25:211-6	IF = 1.475	
106. Childs Nerv Syst 2009;25:111-8	IF = 1.475	
107. Childs Nerv Syst 2009;25:361-365	IF = 1.475	
108. Acta Paediatrica 2009;256:176-182	IF = 1.634	
109. Epilepsy Res 2009;85:89-95	IF = 2.740	
110. Pediatric Neurol 2009; 40:383-386	IF = 3.372	
111. Acta Paediatrica 2009;98:1130-6	IF = 1.634	
112. Acta Paediatrica 2009;99:7	IF = 1.634	
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114. Neuropediatrics 2009;40:186-188	IF = 1.947	
115. Minerva Pediatr 2009;61:557-9	IF = 1.312	[22.478]
116. Eur J Paediatr Neurol 2010;14:192-193	IF = 3.140	
117. Eur J Pediatr 2010;169:475-481	IF = 2.335	
118. Child Nerv Syst 2010;26:133-136	IF = 1.475	
119. Acta Paediatrica 2010;99:460-63	IF = 1.634	
120. Childs Nerv Syst 2010;26:995-1002	IF = 1.475	
121. J Neuroimmunol 2010;223:92-99	IF = 3.178	
122. Pediatr Neurol 2010;43:395-402	IF = 3.372	
123. Neuropediatrics 2010;41:60-65	IF = 1.947	
124. Neuropediatrics 2010;41:246-55	IF = 1.947	
125. Curr Neuroparmacol 2010;8:135-48	IF = 7.363	
126. Front Biosci (Elite Ed) 2010;2:701-10	IF = 2.250	
127. Dev Med Child Neurol 2010;52:700-707	IF = 5.449	[36.371]
128. Area Pediatrica 2010;11:I-XXIII		
129. Childs Nerv Syst 2011;27:635-38	IF = 1.475	
130. Acta Paediatrica 2011;100:121-127	IF = 1.634	
131. Am J Med Genet 2011;155:582-5	IF = 2.802	
132. Pediatr Int 2011;53:964-7	IF = 0.939	
133. Acta Med Medit 2011;27:149-152	IF = 0.219	
134. Acta Med Medit 2011;27:153-162	IF = 0.219	
135. Acta Med Medit 2011;27:163-168	IF = 0.219	
136. Childs Nerv Syst 2011;27:365-71	IF = 1.475	[8.982]

137. Childs Nerv Syst 2012;28:141-5	IF = 1.475	
138. Brain Dev 2012;54:143-147	IF = 1.961	
139. J Hyperten 2012;30:629-630	IF = 4.884	
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141. J Child Neurol 2012;27:657-662	IF = 1.987	
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143. Brain Dev 2012;34:459-468	IF = 1.961	
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145. The Child 2012;1:e-21		
146. The Child 2012;1:e-7		
147. The Child 2012;1:e-9		
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149. Neurol Sci 2012;33:1401-1405	IF = 3.181	
150. Neuro Endocrinol Lett 2012;33:569-573	IF = 0.765	
151. Brain Dev 2012;34:459-468	IF = 1.961	[32.319]
152. Neurocase 2013;19:458-461	IF = 0.883	
153. Eur J Pediatr Neurol 2013;17:97-101	IF = 3.140	
154. J Pediatr 2013;162:217	IF = 4.406	
155. Ophthal Genet 2013;34:178-179	IF = 1.300	
156. Pediatr Neurol 2013;48:73-75	IF = 3.372	
157. Ital J Pediatr 2013;39:3	IF = 2.683	
158. Neurogenetics 2013;14:89-98	IF = 3.860	
159. Neuropediatrics 2013;44:239-244	IF = 1.947	
160. J Pediatr 2013;162:1084	IF = 4.406	
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163. The Child 2013;1(1):e-12		
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165. J Endocrinol Invest 2013;36:1128	IF = 2.660	
166. J Child Neurol 2013;28:1673-76	IF = 1.987	[32.513]
167. Childs Nerv Syst 2014;30:319-25	IF = 1.475	
168. J Clin Neurosci 2014;21:328-330	IF = 1.961	
169. J Child Neurol 2014;29:58-61	IF = 1.987	
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171. Minerva Pediatr 2014;66:17-22.	IF = 1.312	
172. J Pediatr Endocrinol Metab 2014;27:107-115	IF = 1.420	
173. Am J Med Genet 2014;164:1734-43	IF = 2.802	
174. Am J Med Genet 2014;164A:1262-1267	IF = 2.802	
175. Headache 2014; 54:1229.	IF = 5.887	
176. Ital J Pediatr 2014; 40:79	IF = 2.683	
177. Clin EEG Neurosci 2014;47:243-246	IF = 1.843	
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179. Int J Endocrinol 2014;2014:282489	IF = 3.050	[32.916]

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181. Semin Pediatr Neurol 2015;22:207-233	IF = 3.420	
182. Semin Pediatr Neurol 2015;22:240-258	IF = 3.420	
183. JIMD Report 2015;15:39-45	IF = 1.980	
184. Nutrients 2015;7:5532-9.	IF = 5.717	
185. Neurol Sci 2015;36:1173-1180	IF = 3.181	
186. Eur J Pediatr 2015;174:557-563	IF = 3.183	
187. J Child Neurol 2015; 30:654-658	IF = 1.961	
188. Ital J Pediatr 2015; 41:55	IF = 2.634	
189. Am J Med Genet A 2015; 167A:242-51	IF = 2.802	
190. J Pediatr Neurol 2015;13:1-2	IF = 0.210	
191. J Pediatr Neurol 2015;13:3-7	IF = 0.210	
192. J Pediatr Neurol 2015;13:11-15	IF = 0.210	
193. Drug Saf Case Rep 2015;2:6	IF = 1.580	[34.971]
194. Intern Emerg Med 2016;11:273-275	IF = 3.397	
195. Eur J Med Genet 2016;59:283-89	IF = 4.246	
196. Eur J Paediatr Neurol 2016;20:483-88	IF = 3.140	
197. Medicine (Baltimore) 2016;95:e2705	IF = 1.889	
198. Clin Dysmorphol 2016;25:121-127	IF = 0.690	
199. Clin EEG Neuroscie 2016;47:243-246	IF = 1.843	
200. J Pediatr Biochem 2016;5:120-130	IF = 0.134	
201. J Pediatr Biochem 2016;5:109-114	IF = 0.134	
202. J Pediatr Neurol 2016;14:82-88	IF = 0.134	
203. J Pediatr Neurol 2016;14:25-30	IF = 0.134	
204. J Pediatr Biochem 2016;6:3-10	IF = 0.134	
205. J Pediatr Biochem 2016;6:11-18	IF = 0.134	
206. J Pediatr Biochem 2016;6:19-24	IF = 0.134	
207. J Pediatr Biochem 2016;6:25-29	IF = 0.134	
208. J Pediatr Biochem 2016;6:30-38	IF = 0.134	
209. J Pediatr Biochem 2016;6:39-45	IF = 0.134	
210. J Pediatr Biochem 2016;6:46-52	IF = 0.134	
211. J Pediatr Biochem 2016;6:53-59	IF = 0.134	
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214. AJMG C Sem Med Genet 2016;172:288-95	IF = 3.908	
215. Ital J Pediatr 2016;42:78	IF = 2.634	
216. J Neurosci Res 2016;94:1488-1498	IF = 4.164	
217. Acta Otorhinolaryngol Ital 2016; 36:345-367	IF = 2.124	
218. Quant Imag Med Surg 2016;6:515-524	IF = 3.837	
219. Quant Imag Med Surg 2016;6:525-534	IF = 3.837	[38.422]
220. Child's Nerv Syst 2017;33:549-560	IF = 1.475	
221. Eur J Med Genet 2017;60:93-99	IF = 4.246	
222. J Neurosci Res 2017;95:1182-1193	IF = 4.164	
223. Ital J Pediatr 2017;43:6	IF = 2.634	
224. Neurol Sci 2017;38:493-499	IF = 3.181	
225. Childs Nerv Syst 2017;33:933-940	IF = 1.475	
226. Medicine (Baltimore) 2017;96:e6814	IF = 1.889	
227. Lancet Neurol 2017;16:417-418	IF = 44.182	
228. J Pediatric Surg Case Report 2017;24:12-16	IF = 0.200	
229. Neurol Sci 2017;38:1723-1725	IF = 3.181	
230. J Pediatr Neurol 2017;15:84-89	IF = 0.210	[66.836]

231. Curr Vasc Pharmacol 2018;16:499-509	IF = 2.710	
232. Am J Med Genet 2018;176A:515-550	IF = 2.802	
233. Curr Drug Saf 2018;13:131-136	IF = 1.360	
234. J Pediatr Genet 2018;7:29-34	IF = 1.890	
235. Neurogenetics 2018;19(2):77-9	IF = 2.660	
236. Childs Nerv Syst 2018;34:1271-1278	IF = 1.475	
237. J Child Neurol 2018;33:487-492.	IF = 1.632	
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245. J Pediatr Neurol 2018;16:255-264	IF = 0.210	
246. J Pediatr Neurol 2018;16:265-275	IF = 0.210	
247. J Pediatr Neurol 2018;16:276-281	IF = 0.210	
248. J Pediatr Neurol 2018;16:282-287	IF = 0.210	
249. J Pediatr Neurol 2018;16:288-296	IF = 0.210	
250. J Pediatr Neurol 2018;16:297-304	IF = 0.210	
251. J Pediatr Neurol 2018;16:305-312	IF = 0.210	
252. J Pediatr Neurol 2018;16:313-318	IF = 0.210	
253. J Pediatr Neurol 2018;16:319-327	IF = 0.210	
254. J Pediatr Neurol 2018;16:328-337	IF = 0.210	
255. J Pediatr Neurol 2018;16:338-346	IF = 0.210	
256. J Pediatr Neurol 2018;16:347-351	IF = 0.210	
257. J Pediatr Neurol 2018;16:352-361	IF = 0.210	
258. J Pediatr Neurol 2018;16:362-368	IF = 0.210	
259. J Pediatr Neurol 2018;16:369-377	IF = 0.210	
260. Neuropathology 2018;38:577-582	IF = 2.100	
261. Mult Scler Other Demyelin Disord 2018;3:2	IF = 0.078	[112.029]
262. Mol Syndromol 2019;9:253-258.	IF = 1.450	
263. Eur J Med Genet 2019;62:47-54	IF = 4.246	
264. Behav Neurol 2019;2019:3683548	IF = 3.342	
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266. Pediatr Rep 2019;11:8184	IF = 0.500	
267. Nat Commun 2019;10:3094.	IF = 14.919	
268. Epilepsy Res 2019;158:106223	IF = 3.336	
269. Ital J Pediatr 2019;45:159	IF = 2.634	
270. J Pediatr Genet 2019;8:205-211	IF = 1.890	
271. Acta Medica Mediterranea 2019;35:3501-3504	IF = 0.219	
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274. Cells 2020;9:1902.	IF = 4.829	
275. Child Nerv Syst 2020;36:2229-2268	IF = 1.475	
276. Child Nerv Syst 2020;36:2229-2268	IF = 1.475	
277. Epilepsy Behav 2020;112:107361.	IF = 2.937	
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279. Pharmaceuticals (Basel) 2020;13:145.	IF = 5.863	
280. Med Hypothesis 2020;44:110041	IF = 1.538	
281. Mol Genet Genomic Med 2020;8:e1461.	IF = 1.960	
282. Acta Biomedica 2020;19:113-117	IF = 1.352	
283. Eur J Med Genet 2020;63:103957.	IF = 4.246	
284. Clin Dysmorphol 2020;29:202-206.	IF = 0.690	
285. J Pediatr Genet 2020;9:177-182.	IF = 1.089	
286. Front Pediatr 2020;8:550.	IF = 3.418	
287. Mol Genet Genomic Med 2020;8:e1109.	IF = 1.960	
288. BMC Med Genet 2020;21:128.	IF = 1.988	
289. J Pediatr Neurol 2020;18:206-209.	IF = 0.210	
290. J Pediatr Neurol 2020;18:220-224.	IF = 0.210	
291. Children (Basel) 2020;8:16.	IF = 2.078	[44.750]
292. Dev Med Child Neurol 2021;63:343-348.	IF = 5.449	
293. Pediatr Neurol 2021;117:4-9.	IF = 3.352	
294. Ital J Pediatr 2021;47:85	IF = 2.634	
295. Am J Med Genet Part C SMG 2021;187:224-234	IF = 7.100	
296. Harm Reduct J 2021;18:31.	IF = 5.450	
297. Genet Med 2021;23:1506-1513.	IF = 10.540	
298. BMC Med Ethics 2021;22:40	IF = 2.455	
299. Acta Biomed 2021;92:e2021211	IF = 1.350	
300. Am J Perinatol 2021;38:1010-1022	IF = 4.545	
301. Childs Nerv Syst 2021;37:3715-3720.	IF = 1.475	
302. Children (Basel) 2021;8:727.	IF = 2.863	
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309. Children (Basel) 2021;8:637.	IF = 2.863	
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311. Int J Mol Sci 2021;22:10064.	IF = 5.923	
312. J Pediatr Neurol 2021;19:127-131	IF = 0.210	
313. J Pediatr Neurol 2021;19:001-006	IF = 0.210	
314. Adv Neonat Care 2021 30 July	IF = 1.968	
315. Am J Perinatol 2021 Oct 19.	IF = 4.545	
316. Maternal Foetal Med 2021 13 December	IF = 1.230	
317. J Pediatr Neurol 2021 e-first	IF = 0.080	
318. J Pediatr Neurol 2021 e.first	IF = 0.080	
319. J Pediatr Neurol 2021 e.first	IF = 0.080	
320. J Pediatr Neurol 2021 e.first	IF = 0.080	
321. Ital J Pediatr 2021 e-first	IF = 2.683	[92.526]

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327. Expert Rev Neurother 2022;22:169-177.	IF = 4.618	
328. Intern Emerg Med 2022;17:887-909.	IF = 3.200	
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330. Ital J Pediatr 2022;48:29.	IF = 2.638	
331. Front Pediatr 2022;10:858945.	IF = 3.420	
332. Neurol Sci 2022;43:3523-3532.	IF = 3.307	
333. Acta Neurol Belg 2022;1-2.	IF = 2.396	
334. Front Pediatr 2022;10:775356.	IF = 3.420	
335. Eur J Paediatr Neuro. 2022 May 10 Online:A4-A5.	IF = 3.692	
336. Dialogues Clin Neurosci 2022;23:3-13.	IF = 5.986	
337. Genet Med 2022;24:1967-1977.	IF = 8.822	
338. Diagnostics (Basel) 2022;12:1463.	IF = 3.340	
339. Neurol Sci 2022;43:3523-3532.	IF = 3.307	
340. Genet Med 2022;24:2194-2203.	IF = 8.822	
341. Medicine (Baltimore) 2022;101:e29413.	IF = 0.773	
342. Sci Rep 2022;12:10273.	IF = 4.996	
343. Ital J Pediatr 2022;48:82.	IF = 2.683	
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347. Acta Neurol Bel 2022 May 2:1-2.	IF = 2.471	
348. Ital J Pediatr 2022;48:199	IF = 2.683	
349. Acta Biomed 2022; 93:e2022328.	IF = 1.352	
350. Children (Basel) 2022; 9:1767.	IF = 2.835	
351. Neurol Sci 2022;43: 6529-6538.	IF = 3.307	
352. Front Neurol 2022; 13:885897.	IF = 3.770	
353. Front Pediatr 2022; 10:950911.	IF = 3.418	
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357. Ital J Pediatr 2023;49:19.	IF = 4.440	
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359. Children (Basel) 2023;10:841.	IF = 2.440	
360. Front Chem 2023;11:1164014.	IF = 5.545	
361. Transl Pediatr 2023;12:292-300.	IF = 4.047	
362. Acta Neurol Belg 2023;123:687-688.	IF = 3.330	
363. Biomedicines 2023;11:1899.	IF = 4.700	
364. ACS Sens 2023;8:4152-4160.	IF = 26.383	
365. Skin Res Technol 2023;29:e13481.	IF = 2.240	
366. Front Pediatr 2023;11:1210272.	IF = 3.569	
367. Clin Exp Pediatr 2023;66:350-356.	IF = 2.850	
368. Front Chem 2023;11:1164014.	IF = 5.500	
369. Biomedicines. 2023;11:3247.	IF = 4.700	
370. Glob Med Genet. 2023;10:234-239.	IF = 1.700	

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372. Infectious Disease Reports 2023;16:13-25	IF = 3.200	
373. Molecular Diagnosis & Therapy 2023;1:1-2	IF = 4.476	
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376. Minerva Pediatrics 2023;June 16	IF = 2.340	
377. Neuropediatrics 2023;54:315-321	IF = 3.450	
378. Therap Hypother Temp Manag 2023;13:26-36	IF = 1.200	
379. Neuropediatrics 2023;54:297-307	IF = 3.450	
380. J Neurol Neurosurg Neuropsychiatr 2023;94:201-210	IF = 11.000	
381. Acta Neurol Belgica 2023;123:1339-1444	IF = 3.330	
382. Neuropediatrics 2023;54:315-321	IF = 3.450	
383. Minerva Pediatrics 2023;June 16	IF = 2.340	[124.384]
384. Neurogenetics 2024;1:1-10	IF = 3.017	
385. Dev Med Child Neurol 2024;1:1-2.	IF = 4.646	
386. Current Pediatric Reviews 2024;20:370-374	IF = 4.460	
387. Pediatric Research 2024	IF = 3.953	
388. CNS & Neurological Disorders-Drug Targets 2024	IF = 0.830	
389. Topics in Magnetic Resonance Imaging 2024;33:e0310	IF = 0.450	
390. J Clin Pharmacol 2024;64:227-239	IF = 2.300	
391. WFUMB Ultrasound Open 2024;1:100033	IF = 4.345	
392. Mol Diagn Ther 2024; 28:129-130	IF = 4.000	
393. Seizure 2024;117:115-225	IF = 3.000	
394. Proceedings 2024;9:51	IF = 0.400	
395. Front Hum Neurosci 2024 March 20 th	IF = 2.900	
396. Mol Diagn Ther 2024; April 6 th	IF = 4.000	
397. J Pediatr Neurol 2024 [e-first online]	IF = 0.200	
398. J Pediatr Neurol 2024 [e-first online]	IF = 0.200	
399. J Pediatr Neurol 2024 [e-first online]	IF = 0.200	
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404. J Pediatr Neurol 2024 [e-first online]	IF = 0.200	
405. J Pediatr Neurol 2024 [e-first online]	IF = 0.200	
406. Front Hum Neuroscience 2024 24 th April	IF = 3.456	
407. Seizure 2024; 18:156-163.	IF = 4.560	
408. Ital J Pediatr 2024; 50:73	IF = 4.200	
409. Infect Dis Report 2024; 16:13-25	IF = 3.450	
410. Genes (Basel) 2024; 15:848.	IF = 2.800	
411. Front Public Health 2024;12: 1412406.	IF = 3.000	
412. Diagnostics (Basel) 2024;14: 1341.	IF = 3.000	
413. Curr Issues Mol Biol 2024;46: 6112-6120.	IF = 3.100	
414. J Clin Med 2024;13: 3311.	IF = 3.303	
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418. <i>Vaccine Res</i> 2024; 13:225-231.	IF = 3.1454	
419. <i>Ital J Pediatr</i> 2024;50:143.	IF = 4.414	
420. <i>The Journal of Pediatric Academy</i> 2024 October 12	IF = 2.345	
421. <i>IEEE Sensors Journal</i> 2024 October 3	IF = 3.300	
422. <i>Pediatrics & Neonatology</i> 2024 22 December	IF = 2.365	
423. <i>Nutrients</i> 2024;16:4395	IF = 5.675	
424. <i>Mol Cell Pediatr</i> 2024 online 1 December	IF = 2.565	
425. <i>Front Chem</i> 2024; 11:1164014	IF = 9.200	
426. <i>Open Medicine</i> 2024;19:20240979.	IF = 1.230	
427. <i>Children (Basel)</i> 2024; 11:1441.	IF = 2.540	
428. <i>Ital J Pediatr</i> 2024; 50:257.	IF = 3.650	
429. <i>J Clin Pharmacol</i> 2024; 64:227-239.	IF = 3.400	
430. <i>Clin Exp Vaccine Res</i> 2024; 13:225-231	IF = 4.565	[136.545]
431. <i>Glob Med Genet</i> 2025; 12:100033	IF = 2.450	
432. <i>Nutrients</i> 2025; 17:379	IF = 5.67	
433. <i>Epilepsy Open</i> 2025; 10:31-39	IF = 4.650	
434. <i>Nutrients</i> 2025; 17:678.	IF = 5.670	
435. <i>Front Pediatr</i> 2025;13:1597001.	IF = 2.330	
436. <i>Sci Rep</i> 2025;15:20347.	IF = 3.900	
437. <i>Children (Basel)</i> 2025;12:797.	IF = 2.300	
438. <i>Clin Exp Pediatr</i> 2025 May 12.	IF = 3.600	
439. <i>J Child Neurol</i> 2025;8830738251336408.	IF = 1.600	
440. <i>Biomedicines</i> 2025;13:907.	IF = 3.900	
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442. <i>Nutrients</i> 2025;17:1381.	IF = 4.800	
443. <i>Medicina (Kaunas)</i> 2025;61:574.	IF = 2.600	
444. <i>Epilepsy Res</i> 2025;214:107557.	IF = 2.000	
445. <i>J Clin Med</i> 2025;14:2234.	IF = 3.303	
446. <i>Mol Cell Pediatr</i> 2025;12:3.	IF = 1.237	
447. <i>Nutrients</i> 2025;17(4):678.	IF = 2.800	
448. <i>Epilepsia Open</i> 2025; 06 August	IF = 4.086	
449. <i>Pediatrics & Neonatology</i> 2025;56:94-101	IF = 2.300	[44.340]

INTERNATIONAL RESEARCH PAPERS [Scopus, WOS, PubMed]

1994 - 2025 (August)

IMPACT FACTOR (IF) ** TOTAL = 1.640

[** n. 383 peer-reviewed appearing on Scopus/WOS/PubMed con IF]

IF = from the Journal Citation Reports, JCR (ISI), 2025 - <https://www.jcrweb.com>

IMPACT FACTOR (IF) AVERAGE = 3.545

Total number of research papers Scopus/WOS = 426

H-index [Scopus; WOS; Researchgate] = 58

Total number of citations [Scopus; Researchgate] = 9.800

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1994 [IF = 6.290]

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IMPACT FACTOR (IF) ** TOTAL = 1.640

[** n. 383 peer-reviewed appearing on Scopus/WOS/PubMed con IF]

IF = from the Journal Citation Reports, JCR (ISI), 2025 - <https://www.jcrweb.com>

IMPACT FACTOR (IF) AVERAGE = 3.545

Total number of research papers Scopus/WOS = 421

H-index [Scopus; WOS; Researchgate] = 58

Total number of citations [Scopus; Researchgate] = 9.800

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- 50) Salpietro V, Savasta S, Vrotti A, Ruggieri M.
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INVITED SPEAKER AT INTERNATIONAL / NATIONAL MEETINGS

INTERNAZIONAL METINGS

- 1) M. Ruggieri. Segmentary Nf1
1st Golgi Club Weekends in Rome: "Neurofibromatosis type 1"
Rome, Italy, 2-3 December 1995
- 2) M. Ruggieri.
"The neurocutaneous syndromes"
Department of Dermatology, University of Buckinghamshire
Amersham, Gran Bretagna, 16 Gennaio 1996
- 3) M. Ruggieri.
"Hypomelanosis of Ito: clinical and cytogenetic aspects"
Department of Cytogenetics, University of Oxford
Oxford, Gran Bretagna, 3 Luglio 1999
- 4) M. Ruggieri. Mosaicism in neurofibromatosis type 1. A review of clinical and genetic studies.
8th European Neurofibromatosis (ENF) Meeting
Ulm, Germany, 23-26 September 1999.
- 5) M. Ruggieri. Neurological complications of neurofibromatosis type 1 in childhood
9th European Neurofibromatosis Meeting
Venice, Italy, 6-8 April 2001
- 6) M. Ruggieri. Diagnostic implications in the different forms of neurofibromatosis
4th International Meeting of the Polish Society of Pediatric Oncology
Warsaw, Poland, 22-25 October 2002
- 7) M. Ruggieri. "The different forms of neurofibromatosis"
Warsaw, Poland, 23 October 2002
- 8) M. Ruggieri. Tumours in neurocutaneous syndromes
3rd Symposium on "Progress in Molecular Diagnosis and Treatment of Genetic Based Pediatric Malignancies. Vol. 7: n. 1-2
Warsaw, Poland, 22-25 May 2003
- 9) M. Ruggieri. Diagnostic Work-up in neurofibromatosis type 1
2nd International workshop on low grade gliomas and NF1
Padua, Italy, 18-19 November 2003
- 10) M. Ruggieri. Mosaicism in the different forms of neurofibromatosis
2nd International workshop on low grade gliomas and NF1
Padua, Italy, 18-19 November 2003
- 11) M. Ruggieri. Neurofibromatosis
5th Congress European Pediatric Neurology Society (EPNS)
Taormina (ME), 22-25 September 2003
- 12) M. Ruggieri. Multiple sclerosis in childhood: the Italian Database for children under 10 years of age
1st Meeting of the International Pediatric Multiple Sclerosis Study Group (IPMSSG).
Montreal (Canada), 3-6 July 2006
- 13) M. Ruggieri. Pediatric Multiple Sclerosis: The Italian Experience (1st Flag Meeting).
2nd Meeting of the International Pediatric Multiple Sclerosis Study Group (IPMSSG).
Vancouver (Canada), 2-5 June 2008
- 14) M. Ruggieri. Multiple sclerosis update: multiple sclerosis in the pediatric age
8th Mediterranean Neuroscience Congress & 13th Etnean Epilepsy Workshop
Catania, 3-5 June 2010
- 15) M. Ruggieri. Role of microRNA in the therapy of familial tumour syndromes (neurocutaneous disorders).
1st Joint meeting of the Italian Society for Neuroscience and the Israelian Society of Neuroscience.
Catania, 18-21 April 2012
- 16) M. Ruggieri. Natural history of multiple sclerosis in the pediatric age: implications for outcome.
10th Summer School of Neuroscience.
Neuroinflammation in CNS disorders: priming a target for new therapies.
Catania, 7-13 July 2012
- 17) M. Ruggieri. Neurocutaneous disorders
10th Mediterranean Neuroscience Congress & 15th Etnean Epilepsy Workshop
Catania, 12-14 December 2012

- 18) M. Ruggieri. New pharmacological approaches to neurocutaneous disorders.
11th Summer School of Neuroscience. From small molecules to biologic therapies.
Catania, 6-12 July 2013
- 19) M. Ruggieri. Autism and pervasive developmental disorders:
13th Summer School of Neuroscience. Cognition.
Catania, 6-12 July 2015
- 20) M. Ruggieri. Neurofibromatosis
20th International Meeting of the European Group on Neurofibromatosis (EGN)
Abano Terme, 10th September 2016
- 21) M. Ruggieri. Epilepsy in the setting of neurocutaneous disorders
1st Joint EPNS International Meeting of neonatal neurology
Catania, 21-23 May 2018
- 22) M. Ruggieri. Epilepsy in the setting of neurocutaneous disorders
2nd Joint EPNS International Meeting of neonatal neurology
Catania, 11-13 April 2019
- 22) M. Ruggieri. The heart and vessels in neurocutaneous disorders
X World Congress of Pediatric Cardiology
Syracuse, 22-27 May 2019
- 23) M. Ruggieri. Phacomatoses
XVII Congress of the European Pediatric Neurology Society [EPNS]
Athens, 18-20 September 2019
- 24) M. Ruggieri. Neurocutaneous disorders: from the persons behind the syndromes to the molecular pathways and targeted therapies
2nd "Robert J. Gorlin Memorial Lecture"
15th American Academy of Oral and Maxillofacial Pathology [AAOMP] Annual Meeting
Salt Lake City, 24 May 2021
- 25) M. Ruggieri. Neurobiology of epileptogenesis
VII International ILAE Course on infantile epilepsy
Bologna, 3-8 October 2023

NATIONAL MEETINGS

- 1) M. Ruggieri. Neurofibromatosi segmentale (Nf5).
2° Congresso Nazionale sulla neurofibromatosi
Parma, 24-25 Maggio 1996
- 2) M. Ruggieri. Sclerosi Multipla in età pediatrica
Meeting di Neuroimmunologia Pediatrica. Aspetti clinici e patogenetici
Catania, 19 Settembre 1997
- 3) M. Ruggieri. Genetica delle maggiori sindromi neurocutanee.
XXIV Congresso Nazionale della Società Italiana di Neurologia Pediatrica (SINP)
Viterbo, 8-10 Ottobre 1998
- 4) M. Ruggieri. Neurofibromatosi tipo 1. Aspetti clinici ed assistenziali
55° Congresso della Società Italiana di Pediatria (SIP)
Bologna, 19-23 Settembre 1999
- 5) M. Ruggieri. Le varie forme di neurofibromatosi
Convegno Nazionale Associazione Neurofibromatosi (ANF)
Roma, 7 Dicembre 1999
- 6) M. Ruggieri. Neurofibromatosi tipo 1 segmentale (Aggiornamento)
XI Congresso Nazionale della Società di Pediatria Preventiva e Sociale (SIPPS)
Catania, 19 Novembre 1999
- 7) M. Ruggieri. Le neuroimmagini nelle malattie e sindromi neurocutanee
Simposio CNR su: "Esplorazione morfofunzionale del sistema nervoso centrale"
Catania, 17 Dicembre 1999
- 8) M. Ruggieri. Le varianti della neurofibromatosi
Simposio su: "Le sindromi neurocutanee"
Catania, 18 Febbraio 2000
- 9) M. Ruggieri. Neurofibromatosi
Congresso nazionale su: "Aggiornamenti in neuropediatria"
Siena, 4-6 Maggio 2000
- 10) M. Ruggieri. Le neurofibromatosi: Clinica, epidemiologia e diagnosi clinica
5° Congresso nazionale "Associazione Nazionale Neuroradiologia Pediatrica"
Firenze, 11-13 Maggio 2000
- 11) M. Ruggieri. Storia delle curiosità mediche nella neurofibromatosi
Corso Nazionale di Aggiornamento sulla "Storia della Pediatria"
Catania, 8 Luglio 2000
- 12) M. Ruggieri. Neurofibromatosi tipo 1: Protocollo assistenziale.
3° Congresso Nazionale sulla Neurofibromatosi
Ancona, 2-3 Dicembre 2000
- 13) M. Ruggieri. Novità in campo terapeutico nelle neurofibromatosi.
Convegno Nazionale Associazione Neurofibromatosi (ANF)
Parma, 22 Aprile 2001
- 14) M. Ruggieri. Cutis tricolor ed altri mosaicismi cutanei.
Convegno nazionale Gruppo di Studio Genetica Clinica (GENCLI) della S.I.P.
Catania, 19-21 Maggio 2001
- 15) M. Ruggieri. Mosaicismi neurocutanei
XXVII Congresso Società Italiana di Neuropediatria (SINP)
Reggio Emilia, 15-18 Novembre 2001
- 16) M. Ruggieri. Sindromi neurocutanee rare
XXVIII Congresso Società Italiana di Neuropediatria (SINP)
Taranto, 3-6 Novembre 2002
- 17) M. Ruggieri. Le neurofibromatosi: forme localizzate.
1° Corso residenziale nazionale di neurologia pediatrica (SINP)
Pozzilli (IS), 10 Maggio 2003
- 18) M. Ruggieri. Le neurofibromatosi
VIII Corso di aggiornamento sulle malattie genetiche come malattie sociali
Chieti, 10-11 Luglio 2003

- 19) M. Ruggieri, A. Spalice. Malformazioni della fossa cranica posteriore
XXVIII Congresso Società Italiana di Neuropediatria (SINP)
Firenze, 3-6 Dicembre 2003
- 20) P. Iannetti, M. Ruggieri Developmental Pediatrics: neurologia pediatrica in Europa
6° Convegno Nazionale Prospettive in Pediatria
Napoli, 19-21 Febbraio 2004
- 21) M. Ruggieri. Le neurofibromatosi: diagnosi, storia naturale e terapia.
Congresso nazionale sulle sindromi neurocutanee
Lucca, 3-6 Marzo 2004
- 22) M. Ruggieri. Neurofibromatosi e sclerosi tuberosa.
2° Corso residenziale nazionale di neurologia pediatrica
Pozzilli (IS), 21-22 Maggio 2004
- 23) P. Iannetti, M. Ruggieri. Trattamento chirurgico dell'epilessia: la diagnostica per immagini.
Apprendimento attivo in Neuropediatria: IV Giornata
Nocera Inferiore (SA), 18 Settembre 2004
- 24) M. Ruggieri. Le neurofibromatosi
XXIV Congresso Nazionale della Società Italiana di Neurologia (SIN)
Genova, 26 Settembre 2004
- 25) M. Ruggieri. Neurofibromatosi tipo 2 in età pediatrica
XXX Congresso Società Italiana di Neuropediatria (SINP)
Catania, 28-30 Ottobre 2004
- 26) M. Ruggieri. Neurofibromatosi tipo 2: aspetti clinici e diagnostici
4° Congresso Nazionale sulle neurofibromatosi
Napoli, 6-8 Novembre 2004
- 27) M. Ruggieri. Quando e come richiedere l'analisi genetica nelle facomatosi
XVII Congresso Nazionale della Società Italiana di Neuropsichiatria Infantile e dell'Adolescenza (SINPIA)
Modena, 6-10 Novembre 2004
- 28) M. Ruggieri. Mosaicismi neurocutanei
XXIV Congresso Nazionale della Società Italiana di Neurologia (SIN)
Cernobbio, 28 Settembre 2005
- 29) M. Ruggieri. Sindromi del nevo epidermico
XXXI Congresso Società Italiana di Neuropediatria (SINP)
Pavia, 27-30 Ottobre 2005
- 30) M. Ruggieri. Sindromi neurocutanee
XXXII Congresso Società Italiana di Neuropediatria (SINP)
Sasso Marconi (BO), 28-31 Ottobre 2006
- 31) M. Ruggieri. Complicanze neurologiche nelle malattie sistemiche
XXXIII Congresso Società Italiana di Neuropediatria (SINP)
Bolzano, 30 Ottobre-1 Novembre 2007
- 32) M. Ruggieri. Strategie terapeutiche nelle sindromi neurocutanee
XX Congresso Nazionale congiunto SISSMI e SINGEPED
Palermo, 14-16 Ottobre 2008
- 33) M. Ruggieri. Patogenesi delle malattie immunomediate del SNC
XXXIV Congresso Società Italiana di Neuropediatria (SINP)
Napoli, 14-16 Novembre 2008
- 34) M. Ruggieri. Sclerosi multipla in età pediatrica: storia naturale
I Convegno Nazionale Gruppo di Studio di Neuroimmunologia Pediatrica
Catania, 6-7 Giugno 2009
- 35) M. Ruggieri. Malformazioni e tumori vascolari con interessamento del sistema nervoso
XXXV Congresso Società Italiana di Neuropediatria (SINP)
L'Aquila, 14-16 Novembre 2009
- 36) M. Ruggieri. Atassia-Telangiectasia: trasferibilità dei sistemi di valutazione dall'adulto bambino.
Workshop Nazionale sull'Atassia-Telangiectasia: dalla ricerca clinica alla ricerca di base.
Napoli, 14-16 Gennaio 2010
- 37) M. Ruggieri. Neurofibromatosi tipo 2: Manifestazioni cliniche in età pediatrica
V Congresso Nazionale sulle Neurofibromatosi.
Genova, 17-18 Aprile 2010

- 38) M. Ruggieri. Neurofibromatosi tipo 1 e 2: Novità terapeutiche
V Congresso Nazionale sulle Neurofibromatosi.
Genova, 17-18 Aprile 2010
- 39) M. Ruggieri. Studio delle molecole micor-RNA e possibili implicazioni terapeutiche nella sclerosi tuberosa.
Congresso Nazionale Associazione Sclerosi Tuberosa: Incontriamoci in Sicilia.
Catania, 10 Luglio 2010
- 40) M. Ruggieri. La diagnosi di sclerosi tuberosa: introduzione clinica
V Congresso Nazionale sulla Sclerosi Tuberosa: Approccio multidisciplinare alla Sclerosi Tuberosa.
Salemi (TP), 1-3 Ottobre 2010
- 41) M. Ruggieri. Manifestazioni cliniche e storia naturale delle neurofibromatosi
Genetica: Cosa c'è di nuovo. Neurofibromatosi, ieri, oggi, domani
Terni, 6 Novembre 2010
- 42) M. Ruggieri. Neurofibromatosi: nuove strategie terapeutiche
14° Convegno nazionale: Patologia immune malattie orfane
Torino 18-21 Gennaio 2011
- 43) M. Ruggieri. Sclerosi multipla in età pediatrica
Mediterranean Neurosciences Association: Sclerosi Multipla – Stato dell'arte e nuovi paradigmi
Catania, 25 Febbraio 2011
- 44) M. Ruggieri. Mosaicismi neurocutanei
XXVII Congresso Nazionale Società Italiana di Genetica Umana (SIGU)
Milano, 13-16 Novembre 2011
- 45) M. Ruggieri. Cute e sistema nervoso
XXXVI Congresso Società Italiana di Neuropediatria (SINP)
Padova, 17-19 Novembre 2011
- 46) M. Ruggieri. Ominidi pre-umani ed umani arcaici: evidenze fossili e malattie in età pediatrica, cultura, giochi e vita sociale
VIII Congresso Nazionale Congiunto Storia della Pediatria SIN-SIP
Roma, 3 Marzo 2012
- 47) Ruggieri M, Piane M. Atassia-telangiectasia: aspetti biologici e clinici.
Corso di aggiornamento della Società Italiana di Genetica Umana
Difetti di riparazione del DNA: meccanismi e patologie.
Milano, 28 Maggio 2012
- 48) Ruggieri M. NF2: forme ad esordio precoce e nuove strategie farmacologiche.
VI Congresso Nazionale sulle Neurofibromatosi
Bergamo, 27-28 Aprile 2012
- 50) Ruggieri M. Malattie rare in Pediatria: Le sindromi neurocutanee
Expert Meeting sulle Malattie Rare
Palermo, 4-5 Ottobre 2012
- 50) M. Ruggieri. Nuove strategie terapeutiche nelle sindromi neurocutanee.
XXII Congresso Nazionale Congiunto SINGEPED a SSMM
Venezia, 3-5 Novembre 2012
- 51) Ruggieri M. Le malformazioni cerebrali congenite
XIII Meeting Nazionale di Genetica, Immunologia e Pediatria Traslazionale.
Messina, 29-30 Novembre 2012
- 52) Ruggieri M. Sclerosi Multipla Pediatrica. Stato dell'arte: Esperienza della Società Italiana di Pediatria
AISM: Sclerosi Multipla Pediatrica. Stato dell'arte e strategie d'intervento.
Roma, 22 Aprile 2013
- 53) Ruggieri M. Forme sindromiche e non sindromiche associate a spasmi infantili
XIV Meeting Nazionale di Genetica, Immunologia e Pediatria Traslazionale.
Messina, 29-30 Novembre 2013
- 54) M. Ruggieri. Epidemiologia, clinica e storia naturale della sclerosi multipla ad esordio in epoca pre-puberale
XXXVIII Congresso Società Italiana di Neuropediatria (SINP)
Palermo, 27-29 Novembre 2014
- 55) Ruggieri M. Storia naturale della sclerosi multipla in età infantile
XV Meeting Nazionale di Genetica, Immunologia e Pediatria Traslazionale.
Messina, 28-29 Novembre 2014

- 56) Ruggieri M. Macchie cutanee aiuto!
IV Incontro "La Genetica per il Pediatra"
Catanzaro, 3-5 Luglio 2015
- 57) Ruggieri M. ADEM e forme correlate
Meeting Nazionale in Pediatria e Medicina dell'Adolescenza
Catanzaro, 21-24 Ottobre 2015
- 58) Ruggieri M. Analisi delle CNV nella sindrome di Lennoux-Gastaut
Giornata Mondiale sulla sindrome di Lennox-Gastaut
Bologna, 31 Ottobre 2015
- 59) Ruggieri M. Terapie biologiche nelle sindromi neurocutanee.
XVI Meeting Nazionale di Genetica, Immunologia e Pediatria Traslazionale.
Messina, 26-28 Novembre 2015
- 60) M. Ruggieri. Aspetti neuropsicologici della SM in età pediatrica
XXXIX Congresso Nazionale Società Italiana di Neuropediatria (SINP)
Roma, 27-29 Novembre 2015
- 61) M. Ruggieri. Sindromi neurocutanee con malformazioni vascolari
XL Congresso Nazionale Società Italiana di Neuropediatria (SINP)
Firenze, 27-29 Novembre 2016
- 62) M. Ruggieri. Nuove forme di malattie immuno-mediate del SNC in età pediatrica
XLI Congresso Nazionale della Società Italiana di Neurologia Pediatrica (SINP)
Matera, 10-12 Novembre 2017
- 63) M. Ruggieri. Nuove strategie terapeutiche nelle sindromi neurocutanee
XLII Congresso Nazionale della Società Italiana di Neurologia Pediatrica (SINP)
Bologna, 11-13 Novembre 2018
- 64) M. Ruggieri. Neurofibromatosi tipo 1: dalla preistoria all'età moderna e oltre.....
7° Congresso Nazionale sulle neurofibromatosi.
Bressanone (BZ), 25 Maggio 2019
- 65) M. Ruggieri. Neurofibromatosi tipo 1.
74° Congresso Nazionale della Società Italiana di Pediatria
Bologna, 28 Maggio – 1 Giugno 2019
- 66) M. Ruggieri. Sviluppo del sistema nervoso nel feto e nel neonato/Malformazioni cerebrali
XXV Congresso Nazionale della Società Italiana di Neonatologia
Catania, 25-27 Settembre 2019
- 67) M. Ruggieri. Le sindromi malformative ed il laboratorio di genetica per il pediatra
XIII Congresso Nazionale FIMP
Paestum, 16-18 Ottobre 2019
- 68) M. Ruggieri. Facomatosi
XLIII Congresso Nazionale della Società Italiana di Neurologia Pediatrica (SINP)
Napoli, 28-30 Novembre 2019
- 69) M. Ruggieri. Sindromi neurocutanee.
75° Congresso Nazionale della Società Italiana di Pediatria
Webinar, 22-26 Maggio 2020
- 70) M. Ruggieri. Nuove tecniche in genetica molecolare in neonatologi
XXVI Congresso Nazionale della Società Italiana di Neonatologia
Webinar, 28-30 Settembre 2020
- 71) M. Ruggieri. La neurobiologia dello sviluppo
XIV Congresso Nazionale FIMP
Milano, 15-17 Ottobre 2020
- 72) M. Ruggieri. Malattie immuno-mediate del sistema nervoso
XLIV Congresso Nazionale della Società Italiana di Neurologia Pediatrica (SINP)
Webinar, 28-30 Novembre 2020
- 73) M. Ruggieri, A. Polizzi. Le sindromi neurocutanee nella storia e nell'arte
XII Congresso Nazionale "DermArt"
Roma, 24-25 Settembre 2021
- 74) M. Ruggieri. La neurogenetica per il pediatra
17° Forum Nazionale dell'Associazione Nazionale Specializzandi in Pediatria (ONSP)
Riccione, 1 Ottobre 2021

- 75) M. Ruggieri. La genetica clinica e di laboratorio per il pediatra
XV Congresso Nazionale FIMP
Baveno, 7-9 Ottobre 2021
- 76) M. Ruggieri. Manifestazioni neurologiche e psichiatriche durante infezione da COVID-19.
76° Congresso Nazionale della Società Italiana di Pediatria (SIP)
Roma - Webinar, 20 Ottobre 2021
- 77) M. Ruggieri, AD. Praticò. Paralisi cerebrale infantile: work-up dalla diagnosi alla riabilitazione.
XLV Congresso Nazionale della Società Italiana di Neurologia Pediatrica (SINP)
Roma, 25-27 Novembre 2021
- 78) M. Ruggieri. Geni, ambiente e adattamento del cervello del bambino nell'evoluzione umana.
77° Congresso Nazionale della Società Italiana di Pediatria (SIP)
Sorrento, 18-25 maggio 2022
- 79) M. Ruggieri. Basi neurobiologiche della differenza di genere.
9° Congresso Nazionale della Società Italiana di Cure Primarie in Pediatria (SICuPP)
Catania, 9-11 giugno 2022
- 80) M. Ruggieri. Sclerosi multipla e malattie degenerative della sostanza bianca cerebrale.
3° Webinar Nazionale dell'Associazione Italiana per la Sclerosi Multipla (AISM)
Milano, 14 ottobre 2022
- 81) M. Ruggieri. Sindromi malformative complesse: segni dismorfici per la diagnosi.
XLVI Congresso Nazionale della Società Italiana di Neurologia Pediatrica (SINP)
Milano, 20-22 ottobre 2022
- 82) M. Ruggieri. L'evoluzione del cervello del bambino tra geni e ambiente. .
5° Workshop nazionale di Neonatologia
Milano, 4-5 novembre 2022
- 83) M. Ruggieri. L'evoluzione del cervello del bambino e l'interazione cervello/microchip: intelligenza artificiale. .
2° Convegno nazionale della Società Italiana di Ricerca Pediatrica (SIRP)
Napoli, 4-6 giugno 2023
- 84) M. Ruggieri. Terapie geniche in neonatologia. .
XXIX Congresso Nazionale della Società Italiana di Neonatologia
Napoli, 4-6 ottobre 2023
- 85) M. Ruggieri. Lezioni dal passato al futuro: la paleoneurologia. .
XVII Congresso Nazionale della Federazione Italiana dei Medici Pediatri (FIMP)
Taormina, 12-15 ottobre 2023
- 86) M. Ruggieri. Convulsioni febbrili.
78° Congresso Nazionale della Società Italiana di Pediatria (SIP)
Torino, 25-28 ottobre 2023
- 87) M. Ruggieri. Neurofibromatosi tipo 1 / Patogenesi delle sindromi miasteniche.
XLVII Congresso Nazionale della Società Italiana di Neurologia Pediatrica (SINP)
Pisa, 30 novembre - 1° dicembre 2023
- 88) M. Ruggieri. La pediatria nell'ateneo catanese tra '800 e '900. .
101 anni dalla Fondazione della UOC di Clinica Pediatrica dell'Università degli Studi di Catania
Catania, 15 dicembre 2023
- 89) M. Ruggieri. Terapie innovative. .
XXX Congresso Nazionale della Società Italiana di Neonatologia
Roma, 4-6 aprile 2024
- 90) M. Ruggieri. La ricerca pediatrica in ambito internazionale e le società scientifiche italiane.
3° Convegno nazionale della Società Italiana di Ricerca Pediatrica (SIRP)
Roma, 4-6 giugno 2024
- 91) M. Ruggieri. Deficit di neurotrasmettitori
79° Congresso Nazionale della Società Italiana di Pediatria (SIP)
Firenze, 25-28 novembre 2024
- 92) M. Ruggieri. Neurobiologia dello sviluppo cerebrale in endocrinologia
XLVIII Congresso Nazionale della Società Italiana di Neurologia Pediatrica (SINP)
Bari, 30 novembre - 1° dicembre 2024
- 93) M. Ruggieri. Storia della neurologia pediatrica attraverso i secoli
XLVIII Congresso Nazionale della Società Italiana di Neurologia Pediatrica (SINP)
Bari, 30 Novembre - 1 dicembre 2024

- 94) M. Ruggieri. Neurofibromatosi tipo 1 [lettura sponsorizzata da Alexion].
79° Congresso Nazionale della Società Italiana di Pediatria (SIP)
Napoli, 25-28 maggio 2025

REGIONAL MEETINGS

- 1) M. Ruggieri. Idrocefalo: aspetti clinici.
VII Meeting Regionale. La Neurologia Pediatrica nei Vari Aspetti: Autismo e idrocefalo congenito
Catania, 20 Giugno 1995
- 2) M. Ruggieri. Aspetti neurologici della Sclerosi Tuberosa.
Meeting sulla Sclerosi Tuberosa
Catania, 10 Ottobre 1995.
- 3) M. Ruggieri. Neurofibromatosi di tipo segmentale.
Convegno sulle neurofibromatosi
Catania, 4 Giugno 1996
- 4) M. Ruggieri. Neurofibromatosi. Aspetti clinici.
Convegno sulle neurofibromatosi e sulla sclerosi tuberosa
Catania, 17 Giugno 1998
- 5) M. Ruggieri. Aspetti clinici ed assistenziali della neurofibromatosi in età pediatrica.
Convegno regionale Associazione Neurofibromatosi (ANF) sulle neurofibromatosi
Catania, 2 Luglio 1999
- 6) M. Ruggieri. Clinica e follow-up delle neurofibromatosi
Congresso regionale: "Le neurofibromatosi nel bambino: realtà e prospettive"
Reggio Calabria, 4 Marzo 2000
- 7) M. Ruggieri. Le sindromi neurocutanee
Convegno Regionale ACP
Ragusa, 27 Gennaio 2001
- 8) M. Ruggieri. Diagnostica clinico-strumentale del bambino ipotonico
2° Convegno Regionale di Neurologia Pediatrica
Catania, 6-7 Dicembre 2001
- 9) M. Ruggieri. Il pediatra e le sindromi neurocutanee
XXX Congresso Regionale della Società Italiana di Pediatria e
V Congresso Regionale della Società Italiana di Neonatologia
Cefalù (ME), 14-15 Dicembre 2001
- 10) M. Ruggieri. Lesioni cutanee quale segno di patologia neurologica
Convegno di Neurologia Pediatrica
Caltagirone, 20 Aprile 2002
- 11) M. Ruggieri. Novità in campo di neurologia pediatrica: sindromi neurocutanee
Meeting regionale ACP
Acireale, 18 Maggio 2002
- 12) M. Ruggieri. Focus su la "Sclerosi Tuberosa"
2° Convegno Regionale di Neurologia Pediatrica
Catania, 6-7 Dicembre 2002
- 13) M. Ruggieri. Le neurofibromatosi
2° Meeting regionale Genetica clinica GENCLI
Messina, 21-22 Novembre 2003
- 14) M. Ruggieri. Diagnosi e storia naturale delle neurofibromatosi
Convegno sulle Genodermatosi
Troina (EN), 2 Ottobre 2004
- 15) M. Ruggieri. L'esame neurologico nel bambino
Meeting regionale ACP
Acireale, 3 Marzo 2005
- 16) M. Ruggieri. Sclerosi multipla in età infantile
Convegno regionale S.I.P.
Ragusa, 18 Aprile 2006
- 17) M. Ruggieri. Mosaicismo neurocutanei
Convegno sulle Genodermatosi
Troina (EN), 10 Ottobre 2007

- 18) M. Ruggieri. Epilessia e malformazioni cerebrali: Classificazione delle malformazioni cerebrali
Incontro L.I.C.E., Sezione Sicilia: Disturbi del movimento ed epilessia
Ragusa, 27-28 Novembre 2009
- 19) M. Ruggieri. Malattie demielinizzanti: ADEM
Congresso Regionale Società Italiana di Neurologia "Le Patologie Disimmuni del Sistema Nervoso"
Cefalù (PA), 18-19 Dicembre 2009
- 20) M. Ruggieri. Sclerosi Multipla nel bambino
Approcci Terapeutici innovativi per la pratica clinica
Catania, 19-20 Febbraio 2010
- 21) M. Ruggieri. Lesioni vasculo-cutanee e sistema nervoso
Convegni regionali PAIDOS. Cute: organo o sistema
Acireale (CT), 20 Marzo 2010
- 22) M. Ruggieri. Ruolo ed efficacia delle immunoglobuline nelle malattie immunomediate del sistema nervoso centrale in età pediatrica
Aggironamenti in tema di Malattie Disimmuni e Trattamento con IVIG
Siracusa, 13 Novembre 2010
- 23) M. Ruggieri. Encefalomielite acuta in età pediatrica.
I Corso Residenziale: Pratica clinica in neurologia pediatrica con focus sui problemi respiratori nelle patologie neuromuscolari
Catania, 2-3 Luglio 2010
- 24) M. Ruggieri. Studi e Registri epidemiologici pediatrici in Sicilia: sclerosi multipla
39° Congresso Regionale SIP – 14° Congresso Regionale SIN – 7° Congresso Regionale SIMEUP
Cefalù (PA), 11-13 Novembre 2010
- 25) M. Ruggieri. ADEM: risultati dello studio regionale sulle malattie immuno-mediate del sistema nervoso
Convegno di Neonatologia, Farmacologia e Pediatria
Catania, 9-11 Dicembre 2010
- 26) Ruggieri M. Atassie acute
Convegni Paidos - SIP
Acireale (CT), 12 Marzo 2011
- 27) Ruggieri M. Esame obiettivo neurologico nel bambino
Percorsi Pediatrici dell'Alcantara
Acireale, 14 Marzo 2015
- 28) Ruggieri M. Nuove terapie biologiche nelle sindromi neurocutanee
Percorsi Pediatrici Etnei
Catania, 21 Aprile 2015
- 29) Ruggieri M. Malattie immunomediate del sistema nervoso
Congresso Regionale congiunto SIP, SIN; SIMEUP, SIAIP e SINP
Palermo, 27-29 Settembre 2016
- 30) Ruggieri M. La violenza sui minori
Congresso Regionale congiunto SIP, SIN; SIMEUP, SIAIP e SINP
Gela, 11-13 Maggio 2017
- 31) Ruggieri M. Il bambino presitorico
Congresso Regionale congiunto SIP, SIN; SIMEUP, SIAIP e SINP
Siracusa, 18-20 Ottobre 2018
- 32) Ruggieri M. Le Scienze omiche
Congresso Regionale congiunto SIP, SIN; SIMEUP, SIAIP e SINP
Cefalù, 3-5 Ottobre 2019
- 33) Ruggieri M. La neurobiologia dello sviluppo
Congresso Regionale congiunto SIP, SIN; SIMEUP, SIAIP e SINP
Webinar, 3-5 Ottobre 2020
- 34) Ruggieri M. Manifestazioni neurologiche e psichiatriche del COVID-19 nei bambini
Congresso Regionale SIP
- 35) Ruggieri M. Migrazioni e deriva genetica nella storia dell'evoluzione umana
Congresso Regionale congiunto SIMEUP e SINP
Acitrezza, 10-11 Dicembre 2021

TEACHING ACTIVITY

University of Catania

- 2003 - 2004 Adjunct professor (24 hours, 2 credits: Pediatrics MED/38)
Degree Course in Neurophysiopathology, Faculty of Medicine & Surgery,
Catania University
- 2004 - 2005 Adjunct professor (24 hours, 2 credits: Pediatrics MED/38)
Degree Course in Neurophysiopathology, Faculty of Medicine & Surgery,
Catania University
- 2005 - 2006 Adjunct Professor (24 hours, 2 credits: Pediatrics MED/38)
Degree Course in Neurophysiopathology, Faculty of Medicine & Surgery,
Catania University
- 2006 - 2007 Adjunct Professor (24 hours, 2 credits: Pediatrics MED/38)
Degree Course in Neurophysiopathology, Faculty of Medicine & Surgery,
Catania University

University of Catania

- 2009 - 2010 Course Holder "Preventive and Social Paediatrics"
(72 hours, 9 CFU: Paediatrics MED/38)
in the following Degree Courses:
EDUCATION AND TRAINING SCIENCES
EDUCATIONAL SCIENCES FOR CHILDREN
- 2010 - 2011 Course Holder of Child Neuropsychiatry
(48 hours, 6 CFU: Child neuropsychiatry MED/39)
in the following Degree Courses:
CHILDCARE EDUCATOR
EDUCATIONAL SCIENCES FOR CHILDREN
- 2011 - 2012 Course Holder "Pediatrics"
(30 hours, 5 CFU: Pediatrics MED/38)
in the following Degree Courses:
PSYCHOLOGICAL SCIENCES AND TECHNIQUES
- 2011 - 2012 Course Holder "Preventive and Social Paediatrics"
(54 hours, 9 CFU: Paediatrics MED/38)
in the following Degree Courses:
EDUCATIONAL SCIENCES FOR CHILDREN
- 2011 - 2012 Course Holder of "Child Neuropsychiatry"
(36 hours, 6 CFU: Child neuropsychiatry MED/39)
In the following Degree Courses:
EDUCATION AND TRAINING SCIENCES
- 2012 - 2013 Course Holder "Pediatrics"
(30 hours, 5 CFU: Pediatrics MED/38)
in the following Degree Courses:
PSYCHOLOGICAL SCIENCES AND TECHNIQUES

2012 - 2013	Course Holder "Child Neuropsychiatry" (42 hours, 7 CFU: Child neuropsychiatry MED/39) in the following Degree Courses: PSYCHOLOGICAL SCIENCES AND TECHNIQUES
2012 – 2013	Course Holder "Preventive and Social Paediatrics" (54 hours, 9 CFU: Paediatrics MED/38) in the following Degree Courses: EDUCATIONAL SCIENCES FOR CHILDREN
2012 - 2013	Course Holder of "Child Neuropsychiatry" (36 hours, 6 CFU: Child neuropsychiatry MED/39) in the following Degree Courses: EDUCATION AND TRAINING SCIENCES
2013 - 2014	Course Holder "Paediatrics" (30 hours, 5 CFU: Pediatrics MED/38) in the following Degree Courses: PSYCHOLOGICAL SCIENCES AND TECHNIQUES
2013 - 2014	Course Holder "Child Neuropsychiatry" (42 hours, 7 CFU: Child neuropsychiatry MED/39) in the following Degree Courses: PSYCHOLOGICAL SCIENCES AND TECHNIQUES
2013 - 2014	Course Holder "Preventive and Social Paediatrics" (54 hours, 9 CFU: Paediatrics MED/38) in the following Degree Courses: EDUCATIONAL SCIENCES FOR CHILDREN
2013 - 2014	Course Holder of "Child Neuropsychiatry" (36 hours, 6 CFU: Child neuropsychiatry MED/39) in the following Degree Courses: EDUCATION AND TRAINING SCIENCES
2013 - 2014	Teaching "Family Paediatrics" (MED/38) (8 hours, 1 CFU: General and Specialist Paediatrics MED/38 - Pole D) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2014 - 2015	Course Holder "Pediatrics" (30 hours, 5 CFU: Pediatrics MED/38) In the following Degree Courses: PSYCHOLOGICAL SCIENCES AND TECHNIQUES
2014 - 2015	Course Holder "Child Neuropsychiatry" (42 hours, 7 CFU: Child neuropsychiatry MED/39) in the following Degree Courses: PSYCHOLOGICAL SCIENCES AND TECHNIQUES
2014 - 2015	Course Holder "Preventive and Social Paediatrics" (54 hours, 9 CFU: Paediatrics MED/38) in the following Degree Courses: EDUCATIONAL SCIENCES FOR CHILDREN

2015 - 2016	Teaching "Paediatrics" (MED/38) (40 hours, 5 CFU: General and Specialist Paediatrics MED/38 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2015 - 2016	Course Holder "Pediatrics" (40 hours, 5 CFU: Pediatrics MED/38) in the following Degree Courses: PSYCHOLOGICAL SCIENCES AND TECHNIQUES
2015 - 2016	Course Holder "Child Neuropsychiatry" (42 hours, 7 CFU: Child neuropsychiatry MED/39) in the following Degree Courses: PSYCHOLOGICAL SCIENCES AND TECHNIQUES
2015 - 2016	Course Holder "Preventive and Social Paediatrics" (54 hours, 9 CFU: Paediatrics MED/38) in the following Degree Courses: EDUCATIONAL SCIENCES FOR CHILDREN
2015 - 2016	Course Holder of "Child Neuropsychiatry" (36 hours, 6 CFU: Child neuropsychiatry MED/39) in the following Degree Courses: EDUCATIONAL SCIENCES FOR CHILDREN
2015 - 2016	Teaching "Family Paediatrics" (MED/38) (8 hours, 1 CFU: General and Specialist Paediatrics MED/38 - Pole D) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2015 - 2016	Teaching Paediatrics (MED/38) (24 hours: 3 CFU) Degree Course in Orthoptics
2016 - 2017	Teaching Holder "Paediatrics" (MED/38) (32 hours, 4 CFU: General and Specialist Paediatrics MED/38 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2016 - 2017	Course Holder "Pediatrics" (36 hours, 6 CFU: Paediatrics MED/38) in the following Degree Courses: PSYCHOLOGICAL SCIENCES AND TECHNIQUES
2016 - 2017	Course Holder "Preventive and Social Paediatrics" (42 hours, 7 CFU: Pediatrics MED/38) in the following Degree Courses: EDUCATIONAL SCIENCES FOR CHILDREN
2016 - 2017	Course Holder of "Child Neuropsychiatry" (36 hours, 6 CFU: Child neuropsychiatry MED/39) in the following Degree Courses: EDUCATION AND TRAINING SCIENCES
2016 - 2017	Teaching Holder Paediatrics (MED/38) (24 hours: 3 CFU) Degree Course in Orthoptics

2017 - 2018	Teaching Holder "Paediatrics" (MED/38) (32 hours, 4 CFU: General and Specialist Paediatrics MED/38 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2017 - 2018	Course Holder "Paediatrics" (36 hours, 6 CFU: Paediatrics MED/38) in the following Degree Courses: PSYCHOLOGICAL SCIENCES AND TECHNIQUES
2017 - 2018	Course Holder "Preventive and Social Paediatrics" (42 hours, 7 CFU: Pediatrics MED/38) in the following Degree Courses: EDUCATIONAL SCIENCES FOR CHILDREN
2017 - 2018	Course Holder of "Developmental Neurology 60 hours, 10 CFU: General and Specialist Paediatrics MED/38) in the following Degree Courses: EDUCATION AND TRAINING SCIENCES
2017 - 2018	Teaching Holder Paediatrics (MED/38) (24 hours: 3 CFU) Degree Course in Orthoptics
2018 - 2019	Teaching Holder "Paediatrics" (MED/38) (32 hours, 4 CFU: General and Specialist Paediatrics MED/38 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2018 - 2019	Course Holder "Paediatrics" (36 hours, 6 CFU: Paediatrics MED/38) in the following Degree Courses: PSYCHOLOGICAL SCIENCES AND TECHNIQUES
2018 - 2019	Course Holder "Preventive and Social Paediatrics" (42 hours, 7 CFU: Pediatrics MED/38) in the following Degree Courses: EDUCATIONAL SCIENCES FOR CHILDREN
2018 - 2019	Course Holder of "Developmental Neurology 60 hours, 10 CFU: General and Specialist Paediatrics MED/38) in the following Degree Courses: EDUCATION AND TRAINING SCIENCES
2019 - 2020	Teaching Holder "Paediatrics" (MED/38) (32 hours, 4 CFU: General and Specialist Paediatrics MED/38 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2020 - 2021	Teaching Holder "Paediatrics" (MED/38) (32 hours, 4 CFU: General and Specialist Paediatrics MED/38 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY

2021 - 2022	Teaching Holder "Paediatrics" (MED/38) (32 hours, 5 CFU: General and Specialist Paediatrics MED/38 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2021 - 2022	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02 - Pole A) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2021 - 2022	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2021 - 2022	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02 - Pole C) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2021 - 2022	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02 - Polo D) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2021 - 2022	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02) in the following degree courses: BACHELOR'S DEGREE IN OBSTETRICS
2021 - 2022	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02) in the following degree courses: BACHELOR'S DEGREE IN MEDICAL RADIOLOGY IMAGING AND RADIOTHERAPY TECHNIQUES [TRMIR]
2022 - 2023	Teaching Holder "Paediatrics" (MED/38) (32 hours, 5 CFU: General and Specialist Paediatrics MED/38 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2022 - 2023	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02) in the following degree courses: BACHELOR'S DEGREE IN OBSTETRICS
2022 - 2023	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02) in the following degree courses: BACHELOR'S DEGREE IN MEDICAL RADIOLOGY IMAGING AND RADIOTHERAPY TECHNIQUES [TRMIR]

2023 - 2024	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02) in the following degree courses: BACHELOR'S DEGREE IN OBSTETRICS
2023 - 2024	Teaching Holder "Paediatrics" (MED/38) (35 hours, 5 CFU: General and Specialist Paediatrics MED/38 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2023 - 2024	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02) in the following degree courses: BACHELOR'S DEGREE IN OBSTETRICS THREE-YEAR DEGREE IN RADIOLOGY AND RADIODIAGNOSTICS TECHNIQUES
2024 - 2025	Teaching Holder "Paediatrics" (MED/38) (35 hours, 5 credits: General and Specialist Pediatrics MED/38 - Single Centre) in the following degree courses: MASTER'S DEGREE IN NURSING SCIENCES
2024 - 2025	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02) In the following degree courses: BACHELOR'S DEGREE IN OBSTETRICS
2023 – 2025	Teaching Holder "Paediatrics" (MED/38) (35 hours, 5 CFU: General and Specialist Paediatrics MED/38 - Pole B) in the following degree courses: MASTER'S DEGREE IN MEDICINE AND SURGERY
2024 - 2025	Teaching holder (foster) "History of Medicine" (15 hours, 2 CFU: History of Medicine MED/02) in the following degree courses: BACHELOR'S DEGREE IN OBSTETRICS THREE-YEAR DEGREE IN RADIOLOGY AND RADIODIAGNOSTICS TECHNIQUES

Postgraduate Training Programs / Schools of Specialization

2012 - 2013	Teaching holder "Paediatrics" [1° year] School of Specialization in Plastic Surgery University of Catania
2013 - 2014	Teaching holder "Paediatrics" Graduate School of Medical Genetics University of Catania
2013 - 2014	Teaching holder "Paediatrics" [1° and 3° year] School of Specialization in Plastic Surgery University of Catania
2014 - 2015	Teaching holder "Paediatrics" [1° and 3° year] School of Specialization in Plastic Surgery University of Catania
2014 - 2015	Teaching holder "Pediatrics/Pediatric Neurology" [1° and 3° year] Postgraduate Training School in Paediatrics University of Catania
2014 - 2015	Teaching holder "Paediatrics" Postgraduate Training School of Medical Genetics University of Catania
2015 - 2016	Teaching holder "Paediatrics/Paediatric Neurology" [1° and 3° year] Postgraduate School in Paediatrics University of Catania
2015 - 2016	Teaching holder "Paediatrics" Postgraduate Training School of Medical Genetics University of Catania
2016 - 2017	Teaching holder "Pediatrics/Pediatric Neurology" [3° and 5° year] Postgraduate Training School in Paediatrics University of Catania
2016 - 2017	Teaching holder "Paediatrics" Postgraduate Training School of Medical Genetics University of Catania
2016 - 2017	Teaching holder "Paediatrics" [1° and 3° year] Postgraduate Training School in Plastic Surgery University of Catania
2016 - 2017	Teaching holder "Paediatrics" [1° and 3° year] School of Specialization in Gynecology and Obstetrics University of Catania
2017 - 2018	Teaching holder "Pediatrics/Pediatric Neurology" [3° and 5° year] Postgraduate Training School in Paediatrics University of Catania
2017 - 2018	Teaching holder "Paediatrics" Postgraduate Training School of Medical Genetics University of Catania

2017 – 2018	Teaching holder “Paediatrics” [1° and 3° year] School of Specialization in Plastic Surgery University of Catania
2017 – 2018	Teaching holder “Paediatrics” [1° and 3° year] School of Specialization in Gynecology and Obstetrics University of Catania
2018 – 2019	Teaching holder “Paediatrics/Paediatric Neurology” [3° and 5° year] Postgraduate Training School in Paediatrics University of Catania
2018 – 2019	Teaching holder “Paediatrics” Postgraduate Training School of Medical Genetics University of Catania
2018 – 2019	Teaching holder “Paediatrics” [1° and 3° year] School of Specialization in Plastic Surgery University of Catania
2018 – 2019	Teaching holder “Paediatrics” [1° and 3° year] School of Specialization in Gynecology and Obstetrics University of Catania
2019 – 2020	Teaching holder “Paediatrics/Paediatric Neurology” [3° and 5° year] Postgraduate Training School in Paediatrics University of Catania
2019 – 2020	Teaching holder “Paediatrics” Postgraduate School of Medical Genetics University of Catania
2019 – 2020	Teaching holder “Paediatrics” [1° and 3° year] School of Specialization in Plastic Surgery University of Catania
2020 – 2021	Teaching holder “Paediatrics/Paediatric Neurology” [3° and 5° year] Postgraduate Training School in Paediatrics University of Catania
2020 – 2021	Teaching holder “Paediatrics” Postgraduate Training School of Medical Genetics University of Catania
2020 – 2021	Teaching holder “Paediatrics” [1° and 3° year] School of Specialization in Plastic Surgery University of Catania
2020 – 2021	Teaching holder “Paediatrics” [1° and 3° year] School of Specialization in Gynecology and Obstetrics University of Catania

2021 – 2022	Teaching holder “Paediatrics/Paediatric Neurology” [3° and 5° year] Postgraduate School in Paediatrics University of Catania
2021 – 2022	Teaching holder “History of Paediatrics and Paediatric Bioethics” Postgraduate School in Paediatrics University of Catania
2021 – 2022	Teaching holder “Paediatrics” [1° and 3° year] School of Specialization in Plastic Surgery University of Catania
2021 – 2022	Teaching holder “Paediatrics” [1° and 3° year] School of Specialization in Gynecology and Obstetrics University of Catania
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THESIS

<http://www.fmag.unict.it/Public/Uploads/links/Ruggieri.pdf>

Academic year 2009 - 2010

6 Three-year degree thesis, Faculty of Education Sciences

2 Specialist Degree Thesis, Faculty of Education Sciences

Academic year 2010 - 2011

25 Bachelor's Thesis, Faculty of Education

5 Specialist Degree Thesis, Faculty of Education Sciences

Academic year 2011 - 2012

56 Bachelor's Thesis, Department of Educational Sciences

8 Specialist Degree Thesis, Department of Educational Sciences

Academic year 2012 - 2013

66 Bachelor's Thesis, Department of Educational Sciences

9 Specialist Degree Thesis, Department of Educational Sciences

Academic year 2013 - 2014

82 Bachelor's thesis, Department of Educational Sciences

11 Specialist Degree Thesis, Department of Educational Sciences

Academic year 2014 - 2015

86 Bachelor's thesis, Department of Educational Sciences

12 Specialist Degree Thesis, Department of Educational Sciences

Academic year 2015 - 2016

56 Bachelor's Thesis, Department of Educational Sciences

11 Specialist Degree Thesis, Department of Educational Sciences

16 Master's Thesis, School "Faculty of Medicine"

1 Specialization Diploma Thesis in Pediatrics

Academic year 2016 - 2017

25 Bachelor's Thesis, Department of Educational Sciences

4 Specialist Degree Thesis, Department of Educational Sciences

27 Master's Thesis in Medicine and Surgery

2 Specialization Diploma Thesis in Pediatrics

Academic year 2017 - 2018

20 Bachelor's Thesis, Department of Educational Sciences

12 Specialistic Degree Thesis, Department of Educational Sciences

34 Master's Thesis in Medicine and Surgery

2 Specialization Diploma Thesis in Pediatrics

Academic year 2018 - 2019

16 Bachelor's Thesis, Department of Educational Sciences

8 Specialist Degree Thesis, Department of Educational Sciences

39 Master's Thesis in Medicine and Surgery

4 Specialization Diploma Thesis in Pediatrics

Academic year 2019 - 2020

3 Bachelor's Thesis, Department of Educational Sciences

2 Specialist Degree Thesis, Department of Educational Sciences

39 Master's Thesis in Medicine and Surgery

3 Specialization Diploma Thesis in Pediatrics

Academic year 2020 - 2021

- 1 Bachelor's thesis, Department of Educational Sciences
- 2 Specialist Degree Thesis, Department of Educational Sciences
- 27 Master's Thesis in Medicine and Surgery
- 4 Specialization Diploma Thesis in Pediatrics

Academic year 2021 - 2022

- 1 Bachelor's thesis, Department of Educational Sciences
- 2 Specialist Degree Thesis, Department of Educational Sciences
- 27 Master's Thesis in Medicine and Surgery
- 4 Specialization Diploma Thesis in Pediatrics

Academic year 2022 - 2023

- 11 Master's Thesis in Medicine and Surgery
- 4 Specialization Diploma Thesis in Pediatrics

Academic year 2023 - 2024

- 21 Master's Thesis in Medicine and Surgery
- 8 Specialization Diploma Thesis in Pediatrics

Academic year 2024 - 2025

- 23 Master's Thesis in Medicine and Surgery
- 7 Specialization Diploma Thesis in Pediatrics

CLINICAL AND CLINICAL-CARE ACTIVITIES

- Internal student in pediatrics [years 1990 - 1991]

Paediatric Clinic, University of Catania,
clinical care activity in the departments of:
Pediatric oncohematology
clinical care activity at day-hospitals in:
Hemoglobinopathies
Stuffy

- Physician in specialist training in Paediatrics [years 1991 – 1995]

Department of Pediatrics, University of Catania
(supervisor: L. Pavone)
clinical care activity in the departments of:
Neonatology (6 months)
Intensive Care Unit (6 months)
Thalassemia center (4 months)
Infectious diseases (6 months)
General paediatrics (astanteria) (12 months)
Neurology (26 months)

- Clinical Assistant with senior registrar status (GMC rating) [years 1995 – 1999]

contract with the English healthcare system (NHS)
Department of Medical Genetics, Paediatrics, Paediatric Neurology and Neuroradiology, John Radcliffe Hospital,
University of Oxford, Oxford, GB
(S. Supervisors. Huson, M. Pike, A. McShane, P. Anslow)
Periods: 1 June 1995 - 31 May 1996;
1 June 1996 - 31 May 1997;
1 June 1997 - 31 May 1998;
6 July 1998 - 10 September 1998;
15 March 1999 - 26 March 1999
Health Service Provided Abroad recognized by decree of the Ministry of Health no. 0012008-P-DGPROF_02-03-
2020 of March 2, 2020
- From 10 November 1995 to 31 May 1998;
- From 6 July to 10 September 1998
- From 15 March to 26 March 1999

clinical-care activity seconded to the departments of:

General genetic counseling
Syndromology genetic counseling
Von Hippel-Lindau disease
Neurofibromatosis (NF1, NF2, segmental NF)
Tuberous sclerosis
Epileptology
General neurology
Pediatrics

Outpatient manager of neurocutaneous syndromes
Department of Medical Genetics, Churchill Hospital
Oxford Radcliffe Hospital, University of Oxford, UK
November 1996 - March 1999

• Researcher/1° Researcher [years 2000 – 2009]

ISN, CNR, Department of Pediatrics, University of Catania

(Collaboration agreement for clinical/welfare activities CNR-MURST/University of Catania)

Period: 8 August 2000 - 15 November 2009

clinical-care activity seconded to the departments of:

General genetic counseling

Syndromology genetic counseling

Neurocutaneous syndromes

Epileptology

General pediatric neurology

General pediatrics

• Associate Professor of Pediatrics [years 2010-2012]

Agreement between the Faculty of Education - in the person of the Director of the Chair of Preventive and Social Paediatrics, prof. Martino Ruggieri, associate professor of Paediatrics as Coordinator of scientific advice for neurocutaneous diseases (hamartoneoplastic syndromes) - and the AOU "Policlinico-Vittorio Emanuele" of Catania c/o UOC of Neurosurgery, PO "Policlinico - G. Rodolico";

10 August 2010 - 9 February 2010 [Resolution of the AOU "Policlinico-Vittorio Emanuele" of Catania no. 682 of 22 July 2010; agreement with the University of Catania of 10 August 2010];

3 March 2011 - 3 September 2011 [Resolution of the AOU "Policlinico-Vittorio Emanuele" of Catania no. 262 of 2 March 2011; agreement with the University of Catania of 2 March 2011];

11 November 2011 - 11 May 2012 [Resolution of the AOU "Policlinico-Vittorio Emanuele" of Catania no. 1292 of 2 November 2011; agreement with the University of Catania of '11 November 2011]

• Associate Professor of Pediatrics - Head [years 2011-2013]

Pediatric Neurology Clinic & Rare Diseases of the Nervous System, Neurology Unit, Provincial Hospital 2, Hospital S. Elia, Caltanissetta

15 December 2011 - 15 June 2012

16 June 2012 - 15 June 2013

[ASP 2 Caltanissetta/University of Catania resolutions – December 2011, June 2012]

Associate Professor of Pediatrics [years 2014-2015]

Agreement with the Faculty of Education Sciences and with the Director of the Chair of Preventive and Social Paediatrics, prof. Martino Ruggieri, associate professor of Pediatrics as Coordinator of scientific advice for pediatric neurological diseases at the AOU "Policlinico-Vittorio Emanuele" of Catania c/o UOC of Pediatrics and pediatric PS, PO "Vittorio Emanuele" of Catania according to the Collaboration Agreement between the University of Catania and the AOU "Policlinico-Vittorio Emanuele" of Catania;

16 January 2014 - 15 January 2015

[Resolution of the University of Catania, omission of the Minutes of the Department of Education Science Council Meeting of 16 January 2014];

• Full Professor of General and Specialist Paediatrics [years 2015-2016]

Afferent to the Department of Clinical and Experimental Medicine (MEDCLIN), the School "Faculty of Medicine" and the UOC of Pediatrics and PS. Paediatric of the AOU "Policlinico-Vittorio Emanuele" of Catania as Associate Professor of Paediatrics [Rectorial Decree no. 5397/II/13 of 16 January 2015] and as Full Professor of General and Specialist Paediatrics [Rectorial Decree no. 164393 of 29 December 2015] at the AOU "Policlinico-Vittorio Emanuele", PO "Policlinico-G. Rodolico";

[Rectorial Decree no. 164393 of 29 December 2015] at the AOU "Policlinico-Vittorio Emanuele", PO "Policlinico-G. Rodolico";

16 January 2015 - 18 September 2016

Full Professor of General and Specialist Pediatrics - Director [years 2016-to date 2022]
 Insertion in a care regime at the NHS and assignment, as Full Professor of General and Specialist Pediatrics, of the Infradepartmental Program of "Diagnosis, Care Paths and Therapies of Rare Diseases of the Nervous System in Pediatric Age" [Resolution of the DG of the AOU "Policlinico-Vittorio Emanuele" of Catania, no. 1555 of 19 September 2016], and of the Coordination of the "Regional Reference Center for Rare Diseases of the Central and Peripheral Childhood Nervous System" [Resolution of the Department of Health of the Sicilian Region, Official Gazette of the Sicilian Region GURS year 72, no. 12, Friday 16 March 2018, Decree of 28 February 2018 pursuant to the DCPM of 12 January 2017] and at the UOC department of Pediatrics and PS. Paediatric of the AOU "Policlinico-Vittorio Emanuele" of Catania and therefore of the "University Hospital "Policlinico G. Rodolico-San Marco", PO "San Marco" of Catania;
 19 September 2016 - 31 December 2016 [AOUP Report Activities and Results of 3 March 2017]
 1 January 2017 - 31 December 2017 [AOUP Report Activities and Results of 19 February 2018]
 1 January 2018 - 31 December 2018 [AOUP Report Activities and Results of 10 April 2019]
 1 January 2019 - 31 December 2019 [AOUP Report Activities and Results of 30 June 2020]
 1 January 2020 - 31 December 2020 [AOUP Report Activities and Results of 18 March 2021]
 1 January 2021 - 31 December 2021 [AOUP Report Activities and Results of 14 January 2022]
 1 January 2022 - 31 October 2022

November 1°, 2022 - up to date

- Full Professor of General and Specialist Pediatrics - Director [years 2022-to date]
 Placement in a care regime at the NHS and assignment, as Director of the UOC of the Directorate of the Pediatric Clinic UOC, [Resolution of the DG of the AOU "Policlinico" of Catania, no. 1555 of 1 November 2022], and of the Coordination of the "Regional Reference Centers for Rare Diseases of the Central and Peripheral Childhood Nervous System and for Childhood and Adult Hereditary Metabolic Diseases" [Resolution of the Department of Health of the Sicilian Region, of 15 November 2022 and of the AOU "Policlinico" of Catania]

Composition of the Pediatric Clinic UOC [year 2024]

Stays [Building 3, floor 1]

UOS of Extended Neonatal Screening and Hereditary Metabolic Diseases [Building 9, floors 1-.3]

UOS of Child Neuropsychiatry [Building 2, floor 1]

Pediatric Diabetology and Endocrinology CRR πBuilding 2, floor 1]

CRR for Hereditary Metabolic Diseases of Childhood and Adolescence [Building 9, plan 1]

CRR for Rare Diseases of the Central and Peripheral Nervous System in Pediatric Age [Building 3, floor 1]

The undersigned, pursuant to articles 46 and 47 of Presidential Decree no. 445/2000, aware of the criminal sanctions provided for by art. 76 Presidential Decree n. 445/2000, in the case of false declarations, falsehoods in documents, use or exhibition of false documents or documents containing data no longer corresponding to the truth, declares that the above reported corresponds to the truth.

I also declare that the titles and attachments (possibly present to integrate the CV) are, upon request, available in a photostatic copy compliant with the originals.

I authorize the processing of my personal data pursuant to art. 13 legislative decree 30 June 2003 n°196 – "Code regarding the protection of personal data" and art. 13 GDPR 679/16 – "European Regulation

Catania, 18th July 2025

Prof. **Martino Ruggieri**



F.to **Martino Michele Lucio Giovanni RUGGIERI**