

# Marcia G Ribeiro

## List of Publications by Year in descending order

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Version: 2024-02-01

39  
papers

812  
citations

566801

15  
h-index

500791

28  
g-index

42  
all docs

42  
docs citations

42  
times ranked

1260  
citing authors

#	ARTICLE	IF	CITATIONS
1	Sanfilippo syndrome type B: Analysis of patients diagnosed by the <scp>MPS</scp> Brazil Network. American Journal of Medical Genetics, Part A, 2022, 188, 760-767.	0.7	3
2	Agreement Between Alberta Infant Motor Scale Assessment and Maternal Perception of Motor Development in Full-Term Infants. Developmental Neurorehabilitation, 2022, , 1-9.	0.5	0
3	Xâ€linked hypophosphatemic rickets: Description of seven new variants in patients followed up in reference hospitals in Rio de Janeiro. Molecular Genetics & Genomic Medicine, 2022, , e1941.	0.6	2
4	Clinical oral profile of pediatric patients with atopic dermatitis: A crossâ€sectional study. Oral Diseases, 2021, 27, 1834-1846.	1.5	3
5	Association between cytogenetic alteration and the audiometric profile of individuals with Turner syndrome. Brazilian Journal of Otorhinolaryngology, 2021, 87, 728-732.	0.4	3
6	PercepÃ§Ã£o de responsÃ¡veis sobre o envolvimento de crianÃ§as e adolescentes diabÃ©ticos em ocupaÃ§Ãµes durante o distanciamento social devido Ã pandemia por covid-19/ Carers perception on the involvement of diabetic children and adolescents in occupations during the social distancing due to the Covid-19 pandemic. Revista Interinstitucional Brasileira De Terapia Ocupacional, 2021, 5, 352-368.	0.1	0
7	High incidence of acquiring methicillin-resistant Staphylococcus aureus in Brazilian children with Atopic Dermatitis and associated risk factors. Journal of Microbiology, Immunology and Infection, 2020, 53, 724-730.	1.5	13
8	Can simple and low-cost motor function assessments help in the diagnostic suspicion of Duchenne muscular dystrophy?. Jornal De Pediatria, 2020, 96, 503-510.	0.9	1
9	Can simple and low-cost motor function assessments help in the diagnostic suspicion of Duchenne muscular dystrophy?. Jornal De Pediatria (VersÃ£o Em PortuguÃs), 2020, 96, 503-510.	0.2	0
10	Oral mycobiome identification in atopic dermatitis, leukemia, and HIV patients â€“ a systematic review. Journal of Oral Microbiology, 2020, 12, 1807179.	1.2	9
11	Determination of Vitamin D Levels in Patients With Neurofibromatosis Type 1 in the Pediatric Age Group. BMC Clinical Pathology, 2020, 13, 2632010X2092893.	0.7	1
12	Understanding the Landscape of X-linked Variants Causing Intellectual Disability in Females Through Extreme X Chromosome Inactivation Skewing. Molecular Neurobiology, 2020, 57, 3671-3684.	1.9	21
13	High-throughput nucleotide sequencing for bacteriome studies in oral squamous cell carcinoma: a systematic review. Oral and Maxillofacial Surgery, 2020, 24, 387-401.	0.6	6
14	BenefÃcios da inserÃ§Ã£o do fisioterapeuta sobre o perfil de prematuros de baixo risco internados em unidade de terapia intensiva. Fisioterapia E Pesquisa, 2019, 26, 51-57.	0.3	2
15	Frequency of GJB2 mutations in patients with nonsyndromic hearing loss from an ethnically characterized Brazilian population. Brazilian Journal of Otorhinolaryngology, 2019, 85, 92-98.	0.4	1
16	Oral Aspects Identified in Atopic Dermatitis Patients: A Literature Review. Open Dentistry Journal, 2018, 12, 424-434.	0.2	9
17	Decompensated Chronic Heart Failure Reduces Plasma L-carnitine. Archives of Medical Research, 2018, 49, 278-281.	1.5	4
18	Timed motor function tests capacity in healthy children. Archives of Disease in Childhood, 2016, 101, 147-151.	1.0	36

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19	Finding <i>FMR1</i> mosaicism in Fragile X syndrome. <i>Expert Review of Molecular Diagnostics</i> , 2016, 16, 501-507.	1.5	29
20	Alopecia Areata-Part III: Prognosis and Treatment. <i>Skinmed</i> , 2016, 14, 361-365.	0.0	1
21	Novel microduplications at Xp11.22 including <i>HUWE1</i> : clinical and molecular insights into these genomic rearrangements associated with intellectual disability. <i>Journal of Human Genetics</i> , 2015, 60, 207-211.	1.1	24
22	Enzyme replacement therapy for Mucopolysaccharidosis Type I among patients followed within the MPS Brazil Network. <i>Genetics and Molecular Biology</i> , 2014, 37, 23-29.	0.6	19
23	Repercussões clínicas e radiológicas do neurofibroma plexiforme na região pálvica. <i>Radiologia Brasileira</i> , 2014, 47, 326-328.	0.3	1
24	A novel in-frame deletion affecting the BAR domain of <i>OPHN1</i> in a family with intellectual disability and hippocampal alterations. <i>European Journal of Human Genetics</i> , 2014, 22, 644-651.	1.4	24
25	Profile of patients assessed for cochlear implants. <i>Brazilian Journal of Otorhinolaryngology</i> , 2014, 80, 305-310.	0.4	6
26	Betaine-homocysteine methyltransferase 742G>A polymorphism and risk of down syndrome offspring in a Brazilian population. <i>Molecular Biology Reports</i> , 2013, 40, 4685-4689.	1.0	12
27	Dental findings and oral health status in patients with mucopolysaccharidosis: a case series. <i>Acta Odontologica Scandinavica</i> , 2013, 71, 157-167.	0.9	18
28	Mental retardation in Duchenne muscular dystrophy. <i>Jornal De Pediatria</i> , 2012, 88, 6-16.	0.9	32
29	A novel nonsense mutation in <i>KDM5C/JARID1C</i> gene causing intellectual disability, short stature and speech delay. <i>Neuroscience Letters</i> , 2011, 498, 67-71.	1.0	57
30	Moderately progressive Ullrich congenital muscular dystrophy. <i>Jornal De Pediatria</i> , 2011, 88, 93-6.	0.9	1
31	Mucopolysaccharidosis I, II, and VI: brief review and guidelines for treatment. <i>Genetics and Molecular Biology</i> , 2010, 33, 589-604.	0.6	150
32	Novel mutations of the <i>BSCL2</i> and <i>AGPAT2</i> genes in 10 families with Berardinelli-Seip congenital generalized lipodystrophy syndrome. <i>Clinical Endocrinology</i> , 2009, 71, 512-517.	1.2	35
33	Mucopolysaccharidoses in Brazil: What happens from birth to biochemical diagnosis?. <i>American Journal of Medical Genetics, Part A</i> , 2008, 146A, 1741-1747.	0.7	63
34	Single median maxillary central incisor: New data and mutation review. <i>Birth Defects Research Part A: Clinical and Molecular Teratology</i> , 2007, 79, 573-580.	1.6	32
35	A clinical study of 77 patients with mucopolysaccharidosis type II. <i>Acta Paediatrica, International Journal of Paediatrics</i> , 2007, 96, 63-70.	0.7	112
36	Insulin resistance in adolescents with Down syndrome: a cross-sectional study. <i>BMC Endocrine Disorders</i> , 2005, 5, 6.	0.9	34

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37	Concurrence of fragile X and Klinefelter syndromes: report of a new case of paternal nondisjunction. <i>Annales De G�n�tologie</i> , 2003, 46, 53-55.	0.4	3
38	Diploid/triploid mosaicism: Further delineation of the phenotype. <i>American Journal of Medical Genetics Part A</i> , 1994, 52, 399-401.	2.4	28
39	Sangue de cord�o umbilical para uso aut�logo ou grupo de pacientes especiais. <i>Revista Brasileira De Hematologia E Hemoterapia</i> , 0, 31, 36-44.	0.7	1