## Sofia Helena Valente Lemos-Marini

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Fat Distribution and Lipid Profile of Young Adults with Congenital Adrenal Hyperplasia Due to 21â€Hydroxylase Enzyme Deficiency. Lipids, 2021, 56, 101-110.	1.7	5
2	Insulin Resistance in Congenital Adrenal Hyperplasia is Compensated for by Reduced Insulin Clearance. Journal of Clinical Endocrinology and Metabolism, 2021, 106, e1574-e1585.	3.6	7
3	Bioelectrical Impedance Phase Angle and Its Determinants in Patients with Classic Congenital Adrenal Hyperplasia. Journal of the American College of Nutrition, 2021, , 1-8.	1.8	1
4	Cardiovascular dysfunction risk in young adults with congenital adrenal hyperplasia caused by 21â€hydroxylase enzyme deficiency. International Journal of Clinical Practice, 2021, 75, e14233.	1.7	4
5	Adult Height in 299 Patients with Turner Syndrome with or without Growth Hormone Therapy: Results and Literature Review. Hormone Research in Paediatrics, 2021, 94, 63-70.	1.8	6
6	Normal bone health in young adults with 21-hydroxylase enzyme deficiency undergoing glucocorticoid replacement therapy. Osteoporosis International, 2021, , 1.	3.1	1
7	The bone densitometry is normal in Turner syndrome prepubertal patients after height age correction. Revista Brasileira De Saude Materno Infantil, 2021, 21, 1129-1134.	0.5	0
8	Why pediatricians need to know the disorders of sex development: experience of 709 cases in a specialized service. Jornal De Pediatria (Versão Em Português), 2020, 96, 607-613.	0.2	0
9	Novel non-classic CYP21A2 variants, including combined alleles, identified in patients with congenital adrenal hyperplasia. Clinical Biochemistry, 2019, 73, 50-56.	1.9	2
10	Systematic review of quality of life in Turner syndrome. Quality of Life Research, 2018, 27, 1985-2006.	3.1	16
11	FSH may be a useful tool to allow early diagnosis of Turner syndrome. BMC Endocrine Disorders, 2018, 18, 8.	2.2	11
12	Prevalence of Testicular Adrenal Rest Tumor and Factors Associated with Its Development in Congenital Adrenal Hyperplasia. Hormone Research in Paediatrics, 2018, 90, 161-168.	1.8	17
13	Functional and Structural Consequences of Nine <i>CYP21A2</i> Mutations Ranging from Very Mild to Severe Effects. International Journal of Endocrinology, 2016, 2016, 1-10.	1.5	8
14	408 Cases of Genital Ambiguity Followed by Single Multidisciplinary Team during 23 Years: Etiologic Diagnosis and Sex of Rearing. International Journal of Endocrinology, 2016, 2016, 1-9.	1.5	23
15	A new compound heterozygosis for inactivating mutations in the glucokinase gene as cause of permanent neonatal diabetes mellitus (PNDM) in double-first cousins. Diabetology and Metabolic Syndrome, 2015, 7, 101.	2.7	12
16	Growth Curves for Girls with Turner Syndrome. BioMed Research International, 2014, 2014, 1-8.	1.9	16
17	Estimation of percent body fat based on anthropometric measurements in children and adolescents with congenital adrenal hyperplasia due to 21-hydroxylase deficiency. Clinical Nutrition, 2013, 32, 45-50.	5.0	0
18	Accuracy of anthropometric measurements in estimating fat mass in individuals with 21-hydroxylase deficiency. Nutrition, 2012, 28, 984-990.	2.4	7

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19	46,XX DSD and Antley-Bixler syndrome due to novel mutations in the cytochrome P450 oxidoreductase gene. Arquivos Brasileiros De Endocrinologia E Metabologia, 2012, 56, 578-585.	1.3	13
20	Turner syndrome and metabolic derangements: Another example of fetal programming. Early Human Development, 2012, 88, 99-102.	1.8	13
21	Spontaneous puberty in girls with early diagnosis of Turner syndrome. Arquivos Brasileiros De Endocrinologia E Metabologia, 2012, 56, 653-657.	1.3	17
22	Metabolic evaluation of young women with congenital adrenal hyperplasia. Arquivos Brasileiros De Endocrinologia E Metabologia, 2011, 55, 646-652.	1.3	5
23	Social skills in women with Turner Syndrome. Scandinavian Journal of Psychology, 2011, 52, 440-447.	1.5	13
24	Growth hormone effect on body composition in Turner syndrome. Endocrine, 2011, 40, 486-491.	2.3	10
25	Novel deletion alleles carrying CYP21A1P/A2chimeric genes in Brazilian patients with 21-hydroxylase deficiency. BMC Medical Genetics, 2010, 11, 104.	2.1	14
26	Structural aspects of the p.P222Q homozygous mutation of HSD3B2 gene in a patient with congenital adrenal hyperplasia. Arquivos Brasileiros De Endocrinologia E Metabologia, 2010, 54, 768-774.	1.3	10
27	Long-term follow-up of an 8-year-old boy with insulinoma as the first manifestation of a familial form of multiple endocrine neoplasia type 1. Arquivos Brasileiros De Endocrinologia E Metabologia, 2010, 54, 754-760.	1.3	18
28	Phenotypic variability in a family with x-linked adrenoleukodystrophy caused by the p.Trp132Ter mutation. Arquivos Brasileiros De Endocrinologia E Metabologia, 2010, 54, 738-743.	1.3	8
29	Cardiovascular and renal anomalies in Turner syndrome. Revista Da Associação Médica Brasileira, 2010, 56, 655-659.	0.7	35
30	Turner syndrome: a pediatric diagnosis frequently made by non-pediatricians. Jornal De Pediatria, 2010, 86, 121-125.	2.0	11
31	Turner's Syndrome and Subclinical Autoimmune Thyroid Disease: A Two-Year Follow-up Study. Journal of Pediatric Endocrinology and Metabolism, 2009, 22, 109-18.	0.9	4
32	Impairment in Anthropometric Parameters and Body Composition in Females with Classical 21-Hydroxylase Deficiency. Journal of Pediatric Endocrinology and Metabolism, 2009, 22, 519-29.	0.9	7
33	Complement 4 phenotypes and genotypes in Brazilian patients with classical 21-hydroxylase deficiency. Clinical and Experimental Immunology, 2009, 155, 182-188.	2.6	6
34	Perfil clÃnico e laboratorial de pacientes pediátricos e adolescentes com diabetes tipo 1. Jornal De Pediatria, 2009, 85, 490-494.	2.0	9
35	Clinical and laboratory profile of pediatric and adolescent patients with type 1 diabetes. Jornal De Pediatria, 2009, 85, 490-4.	2.0	3
36	Apparent mineralocorticoid excess syndrome in a Brazilian boy caused by the homozygous missense mutation p.R186C in the HSD11B2 gene. Arquivos Brasileiros De Endocrinologia E Metabologia, 2008, 52, 1277-1281.	1.3	10

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37	Absence of mutations in Pax6 gene in three cases of Morning Glory syndrome associated with isolated growth hormone deficiency. Arquivos Brasileiros De Endocrinologia E Metabologia, 2008, 52, 1221-1227.	1.3	12
38	Heterozygosis for CYP21A2 mutation considered as 21-hydroxylase deficiency in neonatal screening. Arquivos Brasileiros De Endocrinologia E Metabologia, 2008, 52, 1388-1392.	1.3	5
39	Spontaneous Final Height in Turner's Syndrome in Brazil. Journal of Pediatric Endocrinology and Metabolism, 2007, 20, 1207-14.	0.9	11
40	Serum 21-Deoxycortisol, 17-Hydroxyprogesterone, and 11-Deoxycortisol in Classic Congenital Adrenal Hyperplasia: Clinical and Hormonal Correlations and Identification of Patients with 11β-Hydroxylase Deficiency among a Large Group with Alleged 21-Hydroxylase Deficiency. Journal of Clinical Endocrinology and Metabolism, 2006, 91, 2179-2184.	3.6	61
41	Frequency of 677C -> T and 1298A -> C polymorphisms in the 5,10-methylenetetrahydrofolate reductase (MTHFR) gene in Turner syndrome individuals. Genetics and Molecular Biology, 2006, 29, 41-44.	1.3	8
42	Labioscrotal island flap in feminizing genitoplasty. Journal of Pediatric Surgery, 2004, 39, 1030-1033.	1.6	14
43	Classical congenital adrenal hyperplasia due to 21-hydroxylase deficiency: a cross-sectional study of factors involved in bone mineral density. Journal of Bone and Mineral Metabolism, 2003, 21, 396-401.	2.7	46
44	Validity of the Use of a Few Hand-Wrist Bones for Assessing Bone Age. Journal of Pediatric Endocrinology and Metabolism, 2003, 16, 541-4.	0.9	3
45	Bone mineralization in Turner syndrome: a transverse study of the determinant factors in 58 patients. Journal of Bone and Mineral Metabolism, 2002, 20, 294-297.	2.7	32
46	A importância dos controles domiciliares na redução de internações em portadores de diabetes mellitus do tipo 1. Arquivos Brasileiros De Endocrinologia E Metabologia, 2000, 44, 215-219.	1.3	0
47	Alterações cardiovasculares na SÃndrome de Turner e correlação cariótipo-fenótipo. , 0, , . 		0