Lucas Santana

List of Publications by Year in descending order

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1040056 1199594 20 170 9 12 citations h-index g-index papers 21 21 21 309 all docs docs citations times ranked citing authors

#	Article	IF	CITATIONS
1	Insights from the genetic characterization of central precocious puberty associated with multiple anomalies. Human Reproduction, 2021, 36, 506-518.	0.9	16
2	Phosphodiesterase 2A and 3B variants are associated with primary aldosteronism. Endocrine-Related Cancer, 2021, 28, 1-13.	3.1	17
3	Comparative Analysis of Different International Criteria (ACMG-AMP vs. TENGEN) Applied to Classification of Missense Germline Allelic Variants in Patients With Multiple Endocrine Neoplasia Type 1 or Suspected to this Syndrome. Journal of the Endocrine Society, 2021, 5, A1014-A1014.	0.2	O
4	Comprehensive Analysis of Clinical Features in Index Cases With Multiple Endocrine Neoplasia Type 1 Refine the Risk Rate for Detection of Mutation Distinguishing Negative-Mutation (Phenocopies) and Positive-Mutation Cases. Journal of the Endocrine Society, 2021, 5, A1014-A1015.	0.2	0
5	Comprehensive Genetic Analysis of 128 Candidate Genes in a Cohort With Idiopathic, Severe, or Familial Osteoporosis. Journal of the Endocrine Society, 2020, 4, bvaa148.	0.2	11
6	Targeted massively parallel sequencing for congenital generalized lipodystrophy. Archives of Endocrinology and Metabolism, 2020, 64, 559-566.	0.6	0
7	Targeted sequencing identifies novel variants in common and rare MODY genes. Molecular Genetics & amp; Genomic Medicine, 2019, 7, e962.	1.2	24
8	Evaluation of <i>SHOX</i> defects in the era of nextâ€generation sequencing. Clinical Genetics, 2019, 96, 261-265.	2.0	9
9	Searching for mutations in the HNF1B gene in a Brazilian cohort with renal cysts and hyperglycemia. Archives of Endocrinology and Metabolism, 2019, 63, 250-257.	0.6	6
10	Genetic diagnosis of congenital hypopituitarism by a target gene panel: novel pathogenic variants in GLI2, OTX2 and GHRHR. Endocrine Connections, 2019, 8, 590-595.	1.9	10
11	SUN-034 Genetic Diagnosis of Congenital Isolated or Combined Growth Hormone Deficiency by Massive Parallel Sequencing Using a Target Gene Panel. Journal of the Endocrine Society, 2019, 3, .	0.2	0
12	<i><scp>PDX1</scp></i> â€ <scp>MODY</scp> and dorsal pancreatic agenesis: New phenotype of a rare disease. Clinical Genetics, 2018, 93, 382-386.	2.0	25
13	Homozygous and Heterozygous Nuclear Lamin A p.R582C Mutation: Different Lipodystrophic Phenotypes in the Same Kindred. Frontiers in Endocrinology, 2018, 9, 458.	3.5	13
14	Germline mutation landscape of multiple endocrine neoplasia type 1 using full gene next-generation sequencing. European Journal of Endocrinology, 2018, 179, 391-407.	3.7	14
15	Clinical application of <scp>ACMGâ€AMP</scp> guidelines in <i><scp>HNF1A</scp></i> and <i><scp>GCK</scp></i> variants in a cohort of <scp>MODY</scp> families. Clinical Genetics, 2017, 92, 388-396.	2.0	19
16	Next-generation sequencing in Brazilian MODY patients: a pilot study. Diabetology and Metabolic Syndrome, 2015, 7, .	2.7	0
17	MODY screening: a new center for molecular genetic diagnosis in Brazil. Diabetology and Metabolic Syndrome, 2015, 7, A213.	2.7	0
18	Detection of congenital generalized lipodystrophy mutations by next-generation sequencing: time for a new approach. Diabetology and Metabolic Syndrome, 2015, 7, .	2.7	0

#	Article	IF	CITATIONS
19	Clonagem molecular do oncogene EZH2 de leucemia mieloide crônica e perspectivas terapêuticas. Mundo Da Saude, 2015, 39, 307-315.	0.1	1
20	Pathogenesis of Primary Aldosteronism: Impact on Clinical Outcome. Frontiers in Endocrinology, 0, 13, .	3.5	4