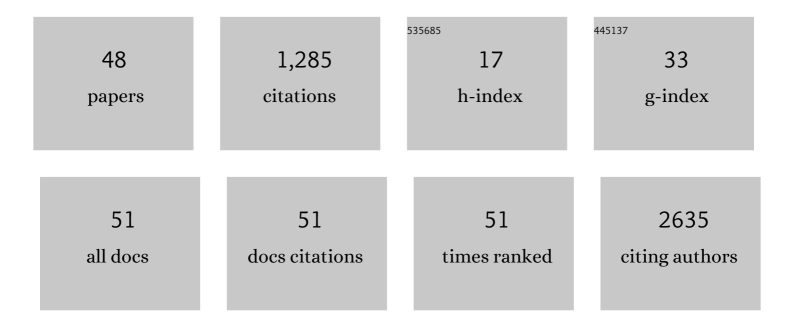
Luisa Azevedo

List of Publications by Year in descending order

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LUISA AZEVEDO

#	Article	lF	CITATIONS
1	Congenital Disorders of Glycosylation in Portugal—Two Decades of Experience. Journal of Pediatrics, 2021, 231, 148-156.	0.9	9
2	SLC35A2-CDG: Novel variant and review. Molecular Genetics and Metabolism Reports, 2021, 26, 100717.	0.4	15
3	Compensatory epistasis explored by molecular dynamics simulations. Human Genetics, 2021, 140, 1329-1342.	1.8	6
4	Common polymorphic <i>OTC</i> variants can act as genetic modifiers of enzymatic activity. Human Mutation, 2021, 42, 978-989.	1.1	6
5	Genetic Variability of the Functional Domains of Chromodomains Helicase DNA-Binding (CHD) Proteins. Genes, 2021, 12, 1827.	1.0	7
6	GBA3: a polymorphic pseudogene in humans that experienced repeated gene loss during mammalian evolution. Scientific Reports, 2020, 10, 11565.	1.6	2
7	The Human Gene Mutation Database (HGMD®): optimizing its use in a clinical diagnostic or research setting. Human Genetics, 2020, 139, 1197-1207.	1.8	353
8	Essential genetic findings in neurodevelopmental disorders. Human Genomics, 2019, 13, 31.	1.4	41
9	Evaluation of InnoQuant® HY and InnoTyper® 21 kits in the DNA analysis of rootless hair samples. Forensic Science International: Genetics, 2019, 39, 61-65.	1.6	10
10	Internal validation of two new retrotransposons-based kits (InnoQuant ® HY and InnoTyper ® 21) at a forensic lab. Forensic Science International, 2018, 283, 1-8.	1.3	7
11	The Yeast Saccharomyces cerevisiae as a Model for Understanding RAS Proteins and their Role in Human Tumorigenesis. Cells, 2018, 7, 14.	1.8	33
12	Improving the in silico assessment of pathogenicity for compensated variants. European Journal of Human Genetics, 2017, 25, 2-7.	1.4	24
13	Major influence of repetitive elements on disease-associated copy number variants (CNVs). Human Genomics, 2016, 10, 30.	1.4	18
14	3-Methylcrotonyl-CoA carboxylase deficiency: Mutational spectrum derived from comprehensive newborn screening. Gene, 2016, 594, 203-210.	1.0	20
15	Trans-species polymorphism in humans and the great apes is generally maintained by balancing selection that modulates the host immune response. Human Genomics, 2015, 9, 21.	1.4	39
16	The mitochondrial genome of the pinewood nematode (Bursaphelenchus xylophilus) lineage introduced in Europe. Mitochondrial DNA, 2014, 25, 420-421.	0.6	2
17	NAMPT and NAPRT1: novel polymorphisms and distribution of variants between normal tissues and tumor samples. Scientific Reports, 2014, 4, 6311.	1.6	21
18	Trimethylaminuria (fish odor syndrome): Genotype characterization among Portuguese patients. Gene, 2013, 527, 366-370.	1.0	16

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19	Novel TTC19 mutation in a family with severe psychiatric manifestations and complex III deficiency. Neurogenetics, 2013, 14, 153-160.	0.7	42
20	Characterization of human NLZ1/ZNF703 identifies conserved domains essential for proper subcellular localization and transcriptional repression. Journal of Cellular Biochemistry, 2013, 114, 120-133.	1.2	18
21	Identification of maternal uniparental isodisomy of chromosome 10 in a patient with mitochondrial DNA depletion syndrome. Molecular Genetics and Metabolism, 2013, 110, 493-494.	0.5	3
22	The Evolutionary Portrait of Metazoan NAD Salvage. PLoS ONE, 2013, 8, e64674.	1.1	8
23	Frequency and Pattern of Heteroplasmy in the Complete Human Mitochondrial Genome. PLoS ONE, 2013, 8, e74636.	1.1	69
24	The Role of Recombination in the Origin and Evolution of Alu Subfamilies. PLoS ONE, 2013, 8, e64884.	1.1	7
25	Characterization of the Human Ornithine Transcarbamylase 3′ Untranslated Regulatory Region. DNA and Cell Biology, 2012, 31, 427-433.	0.9	7
26	Human carbamoyl phosphate synthetase I (CPSI): Insights on the structural role of the unknown function domains. Biochemical and Biophysical Research Communications, 2012, 421, 409-412.	1.0	7
27	Transcriptional regulation of the human mitochondrial peptide deformylase (PDF). Biochemical and Biophysical Research Communications, 2012, 421, 825-831.	1.0	5
28	Successful COG8 and PDF overlap is mediated by alterations in splicing and polyadenylation signals. Human Genetics, 2012, 131, 265-274.	1.8	4
29	Gains, Losses and Changes of Function after Gene Duplication: Study of the Metallothionein Family. PLoS ONE, 2011, 6, e18487.	1.1	67
30	An X-Linked Haplotype of Neandertal Origin Is Present Among All Non-African Populations. Molecular Biology and Evolution, 2011, 28, 1957-1962.	3.5	87
31	Relative frequency of known causes of multiple mtDNA deletions: Two novel POLG mutations. Neuromuscular Disorders, 2011, 21, 483-488.	0.3	16
32	Discussion on common data analysis strategies used in MSâ€based proteomics. Proteomics, 2011, 11, 604-619.	1.3	31
33	Consequences of primer binding-sites polymorphisms on genotyping practice. Open Journal of Genetics, 2011, 01, 15-17.	0.1	9
34	Evolutionary History and Functional Diversification of Phosphomannomutase Genes. Journal of Molecular Evolution, 2010, 71, 119-127.	0.8	11
35	Comparative analyses of the Conserved Oligomeric Golgi (COG) complex in vertebrates. BMC Evolutionary Biology, 2010, 10, 212.	3.2	8
36	Identification of novel L2HGDH gene mutations and update of the pathological spectrum. Journal of Human Genetics, 2010, 55, 55-58.	1.1	6

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37	Ancestral Origin of the ATTCT Repeat Expansion in Spinocerebellar Ataxia Type 10 (SCA10). PLoS ONE, 2009, 4, e4553.	1.1	40
38	Epistatic interactions modulate the evolution of mammalian mitochondrial respiratory complex components. BMC Genomics, 2009, 10, 266.	1.2	33
39	Molecular mechanisms underlying large genomic deletions in ornithine transcarbamylase (<i>OTC</i>) gene. Clinical Genetics, 2009, 75, 457-464.	1.0	24
40	In vitro demonstration of intra-locus compensation using the ornithine transcarbamylase protein as model. Human Molecular Genetics, 2007, 16, 2209-2214.	1.4	15
41	Congenital Disorder of Glycosylation Type Ia: Searching for the Origin of Common Mutations inPMM2. Annals of Human Genetics, 2007, 71, 348-353.	0.3	13
42	Mutational Spectrum and Linkage Disequilibrium Patterns at the Ornithine Transcarbamylase Gene (OTC). Annals of Human Genetics, 2006, 70, 797-801.	0.3	8
43	Epistatic interactions: how strong in disease and evolution?. Trends in Genetics, 2006, 22, 581-585.	2.9	45
44	Novel L2HGDH mutations in 21 patients with L-2-hydroxyglutaric aciduria of Portuguese origin. Human Mutation, 2005, 26, 395-396.	1.1	47
45	Evidence for mutational cis-acting factors affecting mutagenesis in the ornithine transcarbamylase gene. Human Mutation, 2004, 24, 273-273.	1.1	4
46	New polymorphic sites within ornithine transcarbamylase gene: population genetics studies and implications for diagnosis. Molecular Genetics and Metabolism, 2003, 78, 152-157.	0.5	12
47	Ornithine transcarbamylase deficiency: a novel splice site mutation in a family with meiotic recombination and a new useful SNP for diagnosis. Molecular Genetics and Metabolism, 2002, 76, 68-70.	0.5	8
48	Haplotype study of microsatellites flanking the t(15;17) breakpoint in acute promyelocytic leukemia patients from North Portugal. Leukemia, 2002, 16, 1353-1357.	3.3	2