Victor Ef Ferraz

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/7342184/publications.pdf

Version: 2024-02-01

26 papers

482 citations

949033 11 h-index ⁷⁹⁹⁶⁶³
21
g-index

28 all docs

28 docs citations

times ranked

28

884 citing authors

#	Article	IF	CITATIONS
1	Implementação de um programa de mentoring para estudantes de graduação em saúde: a experiência da FMRP-USP. Revista Brasileira De Educacao Medica, 2021, 45, .	0.0	2
2	FMRP unida por um semestre melhor: narrativas de um pacto que celebra valores humanos e compromissos coletivos durante a pandemia da Covid-19. Medicina, 2021, 54, .	0.0	0
3	Lynch syndrome identification in a Brazilian cohort of endometrial cancer screened by a universal approach. Gynecologic Oncology, 2020, 159, 229-238.	0.6	6
4	First description of ultramutated endometrial cancer caused by germline loss-of-function and somatic exonuclease domain mutations in POLE gene. Genetics and Molecular Biology, 2020, 43, e20200100.	0.6	2
5	Molecular basis of familial adenomatous polyposis in the southeast of Brazil: identification of six novel mutations. International Journal of Biological Markers, 2019, 34, 80-89.	0.7	6
6	Perfil de Competência em Genética para Médicos do Brasil: uma Proposta da Sociedade Brasileira de Genética Médica e Genômica. Revista Brasileira De Educacao Medica, 2019, 43, 440-450.	0.0	9
7	Competency Profile in Genetics for Physicians in Brazil: A Proposal of the Brazilian Society of Medical Genetics and Genomics. Revista Brasileira De Educacao Medica, 2019, 43, 440-450.	0.0	2
8	Genetic education, knowledge and experiences between nurses and physicians in primary care in Brazil: A crossâ€sectional study. Australian Journal of Cancer Nursing, 2017, 19, 66-74.	0.8	34
9	Self-reported cancer family history is a useful tool for identification of individuals at risk of hereditary cancer predisposition syndrome at primary care centers in middle-income settings: a longitudinal study. Genetics and Molecular Biology, 2016, 39, 178-183.	0.6	15
10	I Fórum Paulista de Serviços de Apoio ao Estudante de Medicina — Forsa Paulista — "A Carta de MarÃlia― Revista Brasileira De Educacao Medica, 2016, 40, 537-539.	0.0	4
11	The breast cancer immunophenotype of TP53-p.R337H carriers is different from that observed among other pathogenic TP53 mutation carriers. Familial Cancer, 2015, 14, 333-336.	0.9	8
12	Recurrent Mutations in the Basic Domain of TWIST2 Cause Ablepharon Macrostomia and Barber-Say Syndromes. American Journal of Human Genetics, 2015, 97, 99-110.	2.6	61
13	Genotype-Phenotype Correlations in CYP1B1-Associated Primary Congenital Glaucoma Patients Representing Two Large Cohorts from India and Brazil. PLoS ONE, 2015, 10, e0127147.	1.1	18
14	Practicability of comprehensive care in clinical genetics in the brazilian unified health system: expanding the debate. Texto E Contexto Enfermagem, 2014, 23, 1130-1135.	0.4	6
15	Genetics and genomics in Brazil: a promising future. Molecular Genetics & Camp; Genomic Medicine, 2014, 2, 280-291.	0.6	44
16	TP53 p.R337H prevalence in a series of Brazilian hereditary breast cancer families. Hereditary Cancer in Clinical Practice, 2014, 12, 8.	0.6	22
17	Genetic services and testing in Brazil. Journal of Community Genetics, 2013, 4, 355-375.	0.5	66
18	Ablepharon macrostomia syndrome: A distinct genetic entity clinically related to the group of FRAS–FREM complex disorders. American Journal of Medical Genetics, Part A, 2013, 161, 3012-3017.	0.7	7

#	Article	IF	CITATION
19	Predictive models for mutations in mismatch repair genes: implication for genetic counseling in developing countries. BMC Cancer, 2012, 12, 64.	1.1	11
20	Iris coloboma, blepharophimosis, arachnodactyly, joint contractures: Beals syndrome and Van den Ende–Gupta syndrome phenotypic similarities. Clinical Dysmorphology, 2009, 18, 142-144.	0.1	4
21	Mucopolysaccharidoses in Brazil: What happens from birth to biochemical diagnosis?. American Journal of Medical Genetics, Part A, 2008, 146A, 1741-1747.	0.7	63
22	Aggravation of Eyelid and Conjunctival Malignancies Following Photodynamic Therapy in DeSanctis-Cacchione Syndrome. Ophthalmic Plastic and Reconstructive Surgery, 2006, 22, 498-499.	0.4	23
23	Clinical Genetics in Developing Countries: The Case of Brazil. Public Health Genomics, 2004, 7, 95-105.	0.6	23
24	A further case of a Prader-Willi syndrome phenotype in a patient with Angelman syndrome molecular defect. Arquivos De Neuro-Psiquiatria, 2002, 60, 1011-1014.	0.3	8
25	Ablepharon-Macrostomia syndrome: First report of familial occurrence. American Journal of Medical Genetics Part A, 2000, 94, 281-283.	2.4	22
26	Clinical-neurologic, cytogenetic and molecular aspects of the Prader-Willi and Angelman Syndromes. Arquivos De Neuro-Psiquiatria, 1997, 55, 199-208.	0.3	3