Yasser A Al-Sarraj

List of Publications by Year in descending order

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687363 794594 22 427 13 19 citations h-index g-index papers 24 24 24 842 docs citations times ranked citing authors all docs

#	Article	IF	CITATIONS
1	Multiplex epithelium dysfunction due to CLDN10 mutation: the HELIX syndrome. Genetics in Medicine, 2018, 20, 190-201.	2.4	75
2	Whole genome sequencing in the Middle Eastern Qatari population identifies genetic associations with 45 clinically relevant traits. Nature Communications, 2021, 12, 1250.	12.8	37
3	Host Genetic Variants Potentially Associated With SARS-CoV-2: A Multi-Population Analysis. Frontiers in Genetics, 2020, 11, 578523.	2.3	30
4	Qatar genome: Insights on genomics from the Middle East. Human Mutation, 2022, 43, 499-510.	2.5	29
5	T2DM GWAS in the Lebanese population confirms the role of TCF7L2 and CDKAL1 in disease susceptibility. Scientific Reports, 2014, 4, 7351.	3. 3	25
6	CD56 expression in breast cancer induces sensitivity to natural killer-mediated cytotoxicity by enhancing the formation of cytotoxic immunological synapse. Scientific Reports, 2019, 9, 8756.	3.3	23
7	Biallelic <i><scp>SCN</scp>10A</i> mutations in neuromuscular disease and epileptic encephalopathy. Annals of Clinical and Translational Neurology, 2017, 4, 26-35.	3.7	20
8	Type II diabetes mellitus and hyperhomocysteinemia: a complex interaction. Diabetology and Metabolic Syndrome, 2017, 9, 19.	2.7	20
9	A population study of clinically actionable genetic variation affecting drug response from the Middle East. Npj Genomic Medicine, 2022, 7, 10.	3.8	20
10	Thousands of Qatari genomes inform human migration history and improve imputation of Arab haplotypes. Nature Communications, 2021, 12, 5929.	12.8	18
11	Mutations in zinc finger 407 [ZNF407] cause a unique autosomal recessive cognitive impairment syndrome. Orphanet Journal of Rare Diseases, 2014, 9, 80.	2.7	17
12	Multivariate epidemiologic analysis of type 2 diabetes mellitus risks in the Lebanese population. Diabetology and Metabolic Syndrome, 2014, 6, 89.	2.7	17
13	Caffeine Impact on Metabolic Syndrome Components Is Modulated by a CYP1A2 Variant. Annals of Nutrition and Metabolism, 2016, 68, 1-11.	1.9	16
14	Heterozygous PDGFRB Mutation in a Three-generation Family with Autosomal Dominant Infantile Myofibromatosis. Acta Dermato-Venereologica, 2017, 97, 858-859.	1.3	14
15	A chromosomal microdeletion of 15q in a female patient with epilepsy, <scp>ID</scp> , and autism spectrum disorder: a case report. Clinical Case Reports (discontinued), 2017, 5, 1013-1017.	0.5	11
16	Identification of Novel Circulating miRNAs in Patients with Acute Ischemic Stroke. International Journal of Molecular Sciences, 2022, 23, 3387.	4.1	11
17	Distal trisomy 10q syndrome, report of a patient with duplicated q24.31 – qter, autism spectrum disorder and unusual features. Clinical Case Reports (discontinued), 2014, 2, 201-205.	0.5	10
18	Family-Based Genome-Wide Association Study of Autism Spectrum Disorder in Middle Eastern Families. Genes, 2021, 12, 761.	2.4	7

#	Article	IF	CITATIONS
19	Mosaic partial pericentromeric trisomy 8 and maternal uniparental disomy in a male patient with autism spectrum disorder. Clinical Case Reports (discontinued), 2016, 4, 1125-1131.	0.5	4
20	The QChip1 knowledgebase and microarray for precision medicine in Qatar. Npj Genomic Medicine, 2022, $7, 3$.	3.8	4
21	Poking COVID-19: Insights on Genomic Constraints among Immune-Related Genes between Qatari and Italian Populations. Genes, 2021, 12, 1842.	2.4	1
22	The Prevalence and Genetic Spectrum of Familial Hypercholesterolemia in Qatar Based on Whole Genome Sequencing of 14,000 Subjects. Frontiers in Genetics, 0, 13, .	2.3	1