## Zohreh Fattahi

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	SARS oVâ€2 outbreak in Iran: The dynamics of the epidemic and evidence on two independent introductions. Transboundary and Emerging Diseases, 2022, 69, 1375-1386.	1.3	19
2	ZBTB11 dysfunction: spectrum of brain abnormalities, biochemical signature and cellular consequences. Brain, 2022, 145, 2602-2616.	3.7	5
3	Comprehensive <scp>genotypeâ€phenotype</scp> correlation in <scp>AP</scp> â€4 deficiency syndrome; Adding data from a large cohort of Iranian patients. Clinical Genetics, 2021, 99, 187-192.	1.0	2
4	Biallelic mutations in the death domain of PIDD1 impair caspase-2 activation and are associated with intellectual disability. Translational Psychiatry, 2021, 11, 1.	2.4	334
5	<i>CEP104</i> and <i>CEP290</i> ; Genes with Ciliary Functions Cause Intellectual Disability in Multiple Families. Archives of Iranian Medicine, 2021, 24, 364-373.	0.2	3
6	Novel variants in Iranian individuals suspected to have inherited red blood cell disorders, including bone marrow failure syndromes. Haematologica, 2020, 105, e1-e4.	1.7	3
7	Whole genome sequencing identifies a duplicated region encompassing Xq13.2q13.3 in a large Iranian family with intellectual disability. Molecular Genetics & Genomic Medicine, 2020, 8, e1418.	0.6	1
8	Molecular Diagnosis of Hereditary Neuropathies by Whole Exome Sequencing and Expanding the Phenotype Spectrum. Archives of Iranian Medicine, 2020, 23, 426-433.	0.2	6
9	<i>GPR126</i> : A novel candidate gene implicated in autosomal recessive intellectual disability. American Journal of Medical Genetics, Part A, 2019, 179, 13-19.	0.7	12
10	Iranome: A catalog of genomic variations in the Iranian population. Human Mutation, 2019, 40, 1968-1984.	1.1	116
11	Distinct genetic variation and heterogeneity of the Iranian population. PLoS Genetics, 2019, 15, e1008385.	1.5	34
12	Identification of disease ausing variants in the <i>EXOSC</i> gene family underlying autosomal recessive intellectual disability in Iranian families. Clinical Genetics, 2019, 95, 718-725.	1.0	5
13	SZT2 mutation in a boy with intellectual disability, seizures and autistic features. European Journal of Medical Genetics, 2019, 62, 103556.	0.7	12
14	Effect of inbreeding on intellectual disability revisited by trio sequencing. Clinical Genetics, 2019, 95, 151-159.	1.0	49
15	Genetics of intellectual disability in consanguineous families. Molecular Psychiatry, 2019, 24, 1027-1039.	4.1	131
16	Brief Report of Variants Detected in Hereditary Hearing Loss Cases in Iran over a 3-Year Period. Iranian Journal of Public Health, 2019, 48, 1910-1915.	0.3	1
17	<i>CNKSR1</i> gene defect can cause syndromic autosomal recessive intellectual disability. American Journal of Medical Genetics Part B: Neuropsychiatric Genetics, 2018, 177, 691-699.	1.1	4
18	Biallelic missense variants in ZBTB11 can cause intellectual disability in humans. Human Molecular Genetics, 2018, 27, 3177-3188.	1.4	19

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19	De novo Mutation in CACNA1S Gene in a 20-Year-Old Man Diagnosed with Metabolic Myopathy. Archives of Iranian Medicine, 2017, 20, 617-620.	0.2	1
20	Whole Genome Linkage Analysis Followed by Whole Exome Sequencing Identifies Nicastrin (NCSTN) as a Causative Gene in a Multiplex Family with I³-Secretase Spectrum of Autoinflammatory Skin Phenotypes. Journal of Investigative Dermatology, 2016, 136, 1283-1286.	0.3	17
21	Report of limb girdle muscular dystrophy type 2a in 6 Iranian patients, one with a novel deletion in CAPN3 gene. Neuromuscular Disorders, 2016, 26, 277-282.	0.3	5
22	Missense variants in AIMP1 gene are implicated in autosomal recessive intellectual disability without neurodegeneration. European Journal of Human Genetics, 2016, 24, 392-399.	1.4	17
23	Finding mutation within non-coding region of CJB2 reveals its importance in genetic testing of Hearing Loss in Iranian population. International Journal of Pediatric Otorhinolaryngology, 2015, 79, 136-138.	0.4	4
24	A defect in the CLIP1 gene (CLIP-170) can cause autosomal recessive intellectual disability. European Journal of Human Genetics, 2015, 23, 331-336.	1.4	22
25	The Role of a Novel TRMT1 Gene Mutation and Rare GRM1 Gene Defect in Intellectual Disability in Two Azeri Families. PLoS ONE, 2015, 10, e0129631.	1.1	56
26	Report of a patient with limb-girdle muscular dystrophy, ptosis and ophthalmoparesis caused by plectinopathy. Archives of Iranian Medicine, 2015, 18, 60-4.	0.2	8
27	Exome Sequencing and Linkage Analysis Identified Novel Candidate Genes in Recessive Intellectual Disability Associated with Ataxia. Archives of Iranian Medicine, 2015, 18, 670-82.	0.2	4
28	Genetic Investigation of an Iranian Supercentenarian by Whole Exome Sequencing. Archives of Iranian Medicine, 2015, 18, 688-97.	0.2	6
29	Mutation profile of BBS genes in Iranian patients with Bardet–Biedl syndrome: genetic characterization and report of nine novel mutations in five BBS genes. Journal of Human Genetics, 2014, 59, 368-375.	1.1	33
30	Screening for <i>MYO15A</i> gene mutations in autosomal recessive nonsyndromic, <i>GJB2</i> negative Iranian deaf population. American Journal of Medical Genetics, Part A, 2012, 158A, 1857-1864.	0.7	54