## Moshe Frydman

List of Publications by Year in descending order

Source: https://exaly.com/author-pdf/8441038/publications.pdf

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44 papers

4,388 citations

393982 19 h-index 37 g-index

47 all docs

47 docs citations

times ranked

47

4634 citing authors

#	Article	IF	CITATIONS
1	A single ataxia telangiectasia gene with a product similar to PI-3 kinase. Science, 1995, 268, 1749-1753.	6.0	2,634
2	Predominance of null mutations in ataxia-telangiectasia. Human Molecular Genetics, 1996, 5, 433-439.	1.4	247
3	Targeted genomic capture and massively parallel sequencing to identify genes for hereditary hearing loss in middle eastern families. Genome Biology, 2011, 12, R89.	13.9	183
4	High Incidence of Noonan Syndrome Features Including Short Stature and Pulmonic Stenosis in Patients carrying NF1 Missense Mutations Affecting p.Arg1809: Genotype–Phenotype Correlation. Human Mutation, 2015, 36, 1052-1063.	1.1	143
5	Mutations in FYCO1 Cause Autosomal-Recessive Congenital Cataracts. American Journal of Human Genetics, 2011, 88, 827-838.	2.6	132
6	Screening for fragile X syndrome in women of reproductive age. Prenatal Diagnosis, 2000, 20, 611-614.	1.1	117
7	Genetic aspects of Wilson's disease. Journal of Gastroenterology and Hepatology (Australia), 1990, 5, 483-490.	1.4	81
8	Spondyloenchondrodysplasia Due to Mutations in ACP5: A Comprehensive Survey. Journal of Clinical Immunology, 2016, 36, 220-234.	2.0	71
9	A single origin of phenylketonuria in Yemenite Jews. Nature, 1990, 344, 168-170.	13.7	68
10	Velocardiofacial manifestations and microdeletions in schizophrenic inpatients., 1997, 72, 455-461.		68
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		1.1	
11	Noonan syndrome: A cryptic condition in early gestation. , 2000, 92, 159-165.  Consensus interpretation of the p.Met34Thr and p.Val37lle variants in GJB2 by the ClinGen Hearing Loss	1.1	58
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11 12 13	Noonan syndrome: A cryptic condition in early gestation., 2000, 92, 159-165.  Consensus interpretation of the p.Met34Thr and p.Val37lle variants in GJB2 by the ClinGen Hearing Loss Expert Panel. Genetics in Medicine, 2019, 21, 2442-2452.  Novel myosin mutations for hereditary hearing loss revealed by targeted genomic capture and massively parallel sequencing. European Journal of Human Genetics, 2014, 22, 768-775.  Identification of ATM mutations using extended RT-PCR and restriction endonuclease fingerprinting,	1.4	58 56 44
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11 12 13 14	Noonan syndrome: A cryptic condition in early gestation. , 2000, 92, 159-165.  Consensus interpretation of the p.Met34Thr and p.Val37lle variants in GJB2 by the ClinGen Hearing Loss Expert Panel. Genetics in Medicine, 2019, 21, 2442-2452.  Novel myosin mutations for hereditary hearing loss revealed by targeted genomic capture and massively parallel sequencing. European Journal of Human Genetics, 2014, 22, 768-775.  Identification of ATM mutations using extended RT-PCR and restriction endonuclease fingerprinting, and elucidation of the repertoire of A-T mutations in Israel. Human Mutation, 1998, 11, 69-75.  Possible heterogeneity in spondyloenchondrodysplasia: Quadriparesis, basal ganglia calcifications, and chondrocyte inclusions. American Journal of Medical Genetics Part A, 1990, 36, 279-284.  Autosomal recessive Peters anomaly, typical facial appearance, failure to thrive, hydrocephalus, and other anomalies: Further delineation of the Krause-Kivlin syndrome. American Journal of Medical	1.4 1.1 2.4	58 56 44 36 30

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19	Interstitial deletion 2q14q21. American Journal of Medical Genetics Part A, 1989, 34, 476-479.	2.4	28
20	Congenital diaphragmatic hernia in a family segregating a reciprocal translocation $t(5;15)(p15.3;q24)$ ., 2000, 90, 120-122.		23
21	Late onset fulminant Wilson's disease: A case report and review of the literature. World Journal of Gastroenterology, 2014, 20, 17656.	1.4	22
22	Interstitial 7q deletion [46,XY,del (7) (pter â†' cen::q112 â†' qter)] in a retarded quadriplegic boy with normal beta glucuronidase. American Journal of Medical Genetics Part A, 1986, 25, 245-249.	2.4	21
23	MODED: Microcephaly-oculo-digito-esophageal-duodenal syndrome., 1997, 71, 251-257.		21
24	Leukonychia totalis in two sibs. American Journal of Medical Genetics Part A, 1993, 47, 540-541.	2.4	20
25	Familial segregation of cervical ribs, sprengel anomaly, preaxial polydactyly, anal atresia, and urethral obstruction: A new syndrome?. American Journal of Medical Genetics Part A, 1993, 45, 717-720.	2.4	18
26	Váradi syndrome (OFD VI) or opitz trigonocephaly syndrome: Overlapping manifestations in two cousins. American Journal of Medical Genetics Part A, 1993, 47, 451-455.	2.4	18
27	Prenatal diagnosis of a novelCOL1A1 mutation in osteogenesis imperfecta type I carried through full term pregnancy. Prenatal Diagnosis, 2000, 20, 876-880.	1.1	18
28	Oculoâ€palatoâ€cerebral dwarfism: a new syndrome. Clinical Genetics, 1985, 27, 414-419.	1.0	16
29	Spectrum of genes for inherited hearing loss in the Israeli Jewish population, including the novel human deafness gene <scp><i>ATOH1</i></scp> . Clinical Genetics, 2020, 98, 353-364.	1.0	15
30	Alpers progressive infantile neuronal poliodystrophy: An acute neonatal form with findings of the fetal akinesia syndrome. American Journal of Medical Genetics Part A, 1993, 47, 31-36.	2.4	12
31	The Marfan syndrome. Israel Medical Association Journal, 2008, 10, 175-8.	0.1	12
32	Prenatal diagnosis of nail-patella syndrome. , 1999, 19, 287-288.		11
33	Duchenne muscular dystrophy and idiopathic hyperCKemia segregating in a family. American Journal of Medical Genetics Part A, 1995, 58, 209-212.	2.4	10
34	PRENATAL DIAGNOSIS OF RAMBAM–HASHARON SYNDROME. Prenatal Diagnosis, 1996, 16, 266-269.	1.1	9
35	No founder effect detected in Jewish Ashkenazi patients with fragile-X syndrome. Human Genetics, 1997, 101, 186-189.	1.8	8
36	Linkage disequilibrium of common Gaucher disease mutations with a polymorphic site in thepyruvate kinase (PKLR) gene., 1998, 78, 233-236.		8

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37	Refining the Phenotypic Spectrum of KMT5B-Associated Developmental Delay. Frontiers in Pediatrics, 2022, 10, 844845.	0.9	8
38	Ambiguous genitalia in the Proteus syndrome. American Journal of Medical Genetics Part A, 1990, 36, 511-512.	2.4	7
39	t(15;21)(q15;q22.1) pat resulting in partial trisomy and partial monosomy of chromosomes 15 and 21 in two offspring., 1996, 66, 45-51.		6
40	Familial simple hypohidrosis with abnormal palmar dermal ridges. American Journal of Medical Genetics Part A, 1988, 31, 591-596.	2.4	4
41	Syndrome of alopecia totalis and 17b-hydroxysteroid dehydrogenase deficiency. American Journal of Medical Genetics Part A, 1998, 76, 28-31.	2.4	4
42	Autosomalâ€dominant cutis laxa resulting from an intronic mutation in <i><scp>ELN</scp></i> . Experimental Dermatology, 2015, 24, 885-887.	1.4	4
43	Velocardiofacial manifestations and microdeletions in schizophrenic inpatients. American Journal of Medical Genetics Part A, 1997, 72, 455-461.	2.4	3
44	Oligodontia, short stature and small head circumference with normal intelligence. Clinical Genetics, 2008, 46, 316-318.	1.0	2