Manir Ali

List of Publications by Year in descending order

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201575 161767 4,299 66 27 54 citations h-index g-index papers 70 70 70 6229 docs citations citing authors all docs times ranked

#	Article	IF	CITATIONS
1	PDZD8 Disruption Causes Cognitive Impairment in Humans, Mice, and Fruit Flies. Biological Psychiatry, 2022, 92, 323-334.	0.7	14
2	A Recessively Inherited Risk Locus on Chromosome 13q22-31 Conferring Susceptibility to Schizophrenia. Schizophrenia Bulletin, 2021, 47, 796-802.	2.3	3
3	New variants and in silico analyses in GRK1 associated Oguchi disease. Human Mutation, 2021, 42, 164-176.	1.1	7
4	A multi-ethnic genome-wide association study implicates collagen matrix integrity and cell differentiation pathways in keratoconus. Communications Biology, 2021, 4, 266.	2.0	36
5	DYNC2H1 hypomorphic or retina-predominant variants cause nonsyndromic retinal degeneration. Genetics in Medicine, 2020, 22, 2041-2051.	1.1	38
6	Loss-of-Function Mutations in the CFH Gene Affecting Alternatively Encoded Factor H-like 1 Protein Cause Dominant Early-Onset Macular Drusen. Ophthalmology, 2019, 126, 1410-1421.	2.5	25
7	Mutation spectrum and clinical investigation of achromatopsia patients with mutations in the <i>GNAT2</i> gene. Human Mutation, 2019, 40, 1145-1155.	1.1	15
8	Matrix metalloproteinases in keratoconus – Too much of a good thing?. Experimental Eye Research, 2019, 182, 137-143.	1.2	49
9	LHFPL5 mutation: A rare cause of non-syndromic autosomal recessive hearing loss. European Journal of Medical Genetics, 2019, 62, 103592.	0.7	6
10	Biallelic sequence and structural variants in RAX2 are a novel cause for autosomal recessive inherited retinal disease. Genetics in Medicine, 2019, 21, 1319-1329.	1.1	15
11	DETAILED RETINAL IMAGING IN CARRIERS OF OCULAR ALBINISM. Retina, 2018, 38, 620-628.	1.0	13
12	Identification of Inherited Retinal Disease-Associated Genetic Variants in 11 Candidate Genes. Genes, 2018, 9, 21.	1.0	20
13	Association of Genetic Variants With Response to Anti–Vascular Endothelial Growth Factor Therapy in Age-Related Macular Degeneration. JAMA Ophthalmology, 2018, 136, 875.	1.4	30
14	Use of a gene-based case-control association approach in exome sequencing data to elucidate the molecular basis of a mendelian phenotype. Lancet, The, 2017, 389, S14.	6.3	1
15	Defects in the Cell Signaling Mediator \hat{l}^2 -Catenin Cause the Retinal Vascular Condition FEVR. American Journal of Human Genetics, 2017, 100, 960-968.	2.6	74
16	Association of Steroid 5î±-Reductase Type 3 Congenital Disorder of Glycosylation With Early-Onset Retinal Dystrophy. JAMA Ophthalmology, 2017, 135, 339.	1.4	43
17	The effect of <i><scp>COMT</scp> Val158Met</i> and <i><scp>DRD</scp>2 C957T</i> polymorphisms on executive function and the impact of early life stress. Brain and Behavior, 2017, 7, e00695.	1.0	31
18	Specific Alleles of <i>CLN7</i> / <i>MFSD8</i> , a Protein That Localizes to Photoreceptor Synaptic Terminals, Cause a Spectrum of Nonsyndromic Retinal Dystrophy., 2017, 58, 2906.		35

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19	Mutations in the polyglutamylase gene <i>TTLL5</i> , expressed in photoreceptor cells and spermatozoa, are associated with cone-rod degeneration and reduced male fertility. Human Molecular Genetics, 2016, 25, ddw282.	1.4	27
20	Mutations in REEP6 Cause Autosomal-Recessive Retinitis Pigmentosa. American Journal of Human Genetics, 2016, 99, 1305-1315.	2.6	121
21	Homozygous single base deletion in <i>TUSC3</i> causes intellectual disability with developmental delay in an Omani family. American Journal of Medical Genetics, Part A, 2016, 170, 1826-1831.	0.7	14
22	Disease Expression in Autosomal Recessive Retinal Dystrophy Associated With Mutations in the <i>DRAM2 </i>		13
23	Association Between Missense Mutations in the <i>BBS2</i> Gene and Nonsyndromic Retinitis Pigmentosa. JAMA Ophthalmology, 2015, 133, 312.	1.4	43
24	A missense variant in CST3 exerts a recessive effect on susceptibility to age-related macular degeneration resembling its association with Alzheimer's disease. Human Genetics, 2015, 134, 705-715.	1.8	30
25	Biallelic Mutations in the Autophagy Regulator DRAM2 Cause Retinal Dystrophy with Early Macular Involvement. American Journal of Human Genetics, 2015, 96, 948-954.	2.6	42
26	Novel C8orf37 mutations cause retinitis pigmentosa in consanguineous families of Pakistani origin. Molecular Vision, 2015, 21, 236-43.	1.1	10
27	Mutation Screening of Retinal Dystrophy Patients by Targeted Capture from Tagged Pooled DNAs and Next Generation Sequencing. PLoS ONE, 2014, 9, e104281.	1.1	20
28	Spectral domain optical coherence tomography imaging of the posterior segment of the eye in the retinal dysplasia and degeneration chicken, an animal model of inherited retinal degeneration. Veterinary Ophthalmology, 2014, 17, 113-119.	0.6	13
29	Congenital Hereditary Endothelial Dystrophy Caused by SLC4A11 Mutations Progresses to Harboyan Syndrome. Cornea, 2014, 33, 247-251.	0.9	47
30	Simple and Efficient Identification of Rare Recessive Pathologically Important Sequence Variants from Next Generation Exome Sequence Data. Human Mutation, 2013, 34, 945-952.	1.1	4
31	An X-Ray Scattering Study into the Structural Basis of Corneal Refractive Function in an Avian Model. Biophysical Journal, 2013, 104, 2586-2594.	0.2	5
32	Recessive Mutations in SLC38A8 Cause Foveal Hypoplasia and Optic Nerve Misrouting without Albinism. American Journal of Human Genetics, 2013, 93, 1143-1150.	2.6	71
33	Patterns of inheritance, not always easily visible. BMJ, The, 2013, 347, f6610-f6610.	3.0	0
34	Mutational Analysis of <i>MIR184</i> i>in Sporadic Keratoconus and Myopia., 2013, 54, 5266.		73
35	<i>ZNF408</i> is mutated in familial exudative vitreoretinopathy and is crucial for the development of zebrafish retinal vasculature. Proceedings of the National Academy of Sciences of the United States of America, 2013, 110, 9856-9861.	3.3	144
36	Retinal Phenotypes in Patients Homozygous for the G1961E Mutation in the ABCA4Gene., 2013, 54, 520.		3

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37	A new recessively inherited disorder composed of foveal hypoplasia, optic nerve decussation defects and anterior segment dysgenesis maps to chromosome 16q23.3-24.1. Molecular Vision, 2013, 19, 2165-72.	1.1	19
38	Recessive Mutations in <i> TSPAN12 < /i > Cause Retinal Dysplasia and Severe Familial Exudative Vitreoretinopathy (FEVR). , 2012, 53, 2873.</i>		64
39	Next generation sequencing identifies mutations in Atonal homolog 7 (ATOH7) in families with global eye developmental defects. Human Molecular Genetics, 2012, 21, 776-783.	1.4	66
40	<i>CFH</i> , <i>VEGF</i> and <i>HTRA1</i> promoter genotype may influence the response to intravitreal ranibizumab therapy for neovascular age-related macular degeneration. British Journal of Ophthalmology, 2012, 96, 208-212.	2.1	98
41	Rapid Visualisation of Microarray Copy Number Data for the Detection of Structural Variations Linked to a Disease Phenotype. PLoS ONE, 2012, 7, e43466.	1.1	1
42	Profiling Retinal Biochemistry in the MPDZ Mutant Retinal Dysplasia and Degeneration Chick: A Model of Human RP and LCA., 2012, 53, 413.		9
43	Identification of autosomal recessive disease loci using out-bred nuclear families. Human Mutation, 2012, 33, 338-342.	1.1	4
44	Genetic Heterogeneity for Recessively Inherited Congenital Cataract Microcornea with Corneal Opacity., 2011, 52, 4294.		13
45	<i>Mpdz</i> Null Allele in an Avian Model of Retinal Degeneration and Mutations in Human Leber Congenital Amaurosis and Retinitis Pigmentosa. , 2011 , 52 , 7432 .		24
46	Homozygous Mutations in PXDN Cause Congenital Cataract, Corneal Opacity, and Developmental Glaucoma. American Journal of Human Genetics, 2011, 89, 464-473.	2.6	68
47	Changing the status quo bias. British Journal of Ophthalmology, 2011, 95, 1034-1034.	2.1	0
48	The Influence of Lamellar Orientation on Corneal Material Behavior: Biomechanical and Structural Changes in an Avian Corneal Disorder., 2011, 52, 1243.		18
49	Reply to Papanikolaou et al. British Journal of Ophthalmology, 2011, 95, 890-890.	2.1	4
50	The D153del Mutation in GNB3 Gene Causes Tissue Specific Signalling Patterns and an Abnormal Renal Morphology in Rge Chickens. PLoS ONE, 2011, 6, e21156.	1.1	11
51	Genotype-Phenotype Correlation for Leber Congenital Amaurosis in Northern Pakistan. JAMA Ophthalmology, 2010, 128, 107.	2.6	64
52	Mutations in TSPAN12 Cause Autosomal-Dominant Familial Exudative Vitreoretinopathy. American Journal of Human Genetics, 2010, 86, 248-253.	2.6	161
53	Homozygous FOXE3 mutations cause non-syndromic, bilateral, total sclerocornea, aphakia, microphthalmia and optic disc coloboma. Molecular Vision, 2010, 16, 1162-8.	1.1	33
54	Mutations involved in Aicardi-Goutià res syndrome implicate SAMHD1 as regulator of the innate immune response. Nature Genetics, 2009, 41, 829-832.	9.4	610

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55	Null Mutations in LTBP2 Cause Primary Congenital Glaucoma. American Journal of Human Genetics, 2009, 84, 664-671.	2.6	255
56	Loss of the Metalloprotease ADAM9 Leads to Cone-Rod Dystrophy in Humans and Retinal Degeneration in Mice. American Journal of Human Genetics, 2009, 84, 683-691.	2.6	76
57	Ultrastructural changes in the retinopathy, globe enlarged (rge) chick cornea. Journal of Structural Biology, 2009, 166, 195-204.	1.3	33
58	Collagen organization in the chicken cornea and structural alterations in the retinopathy, globe enlarged (rge) phenotype—An X-ray diffraction study. Journal of Structural Biology, 2008, 161, 1-8.	1.3	30
59	A missense mutation in the nuclear localization signal sequence of CERKL (p.R106S) causes autosomal recessive retinal degeneration. Molecular Vision, 2008, 14, 1960-4.	1.1	25
60	Differential Display Reverse Transcription-Polymerase Chain Reaction to Identify Novel Biomolecules in Arthritis Research. Methods in Molecular Medicine, 2007, 136, 329-347.	0.8	0
61	Clinical phenotype associated with homozygosity forÂaÂHOXD13 7-residue polyalanine tract expansion. European Journal of Medical Genetics, 2006, 49, 396-401.	0.7	19
62	Mutations in genes encoding ribonuclease H2 subunits cause Aicardi-Goutià res syndrome and mimic congenital viral brain infection. Nature Genetics, 2006, 38, 910-916.	9.4	592
63	Mutations in the gene encoding the 3′-5′ DNA exonuclease TREX1 cause Aicardi-Goutières syndrome at the AGS1 locus. Nature Genetics, 2006, 38, 917-920.	9.4	752
64	Mutation in the Guanine Nucleotide–Binding Protein β-3 Causes Retinal Degeneration and Embryonic Mortality in Chickens. , 2006, 47, 4714.		41
65	Application of differential display to immunological research. Journal of Immunological Methods, 2001, 250, 29-43.	0.6	23
66	Rheumatoid arthritis synovial T cells regulate transcription of several genes associated with antigen-induced anergy. Journal of Clinical Investigation, 2001, 107, 519-528.	3.9	36