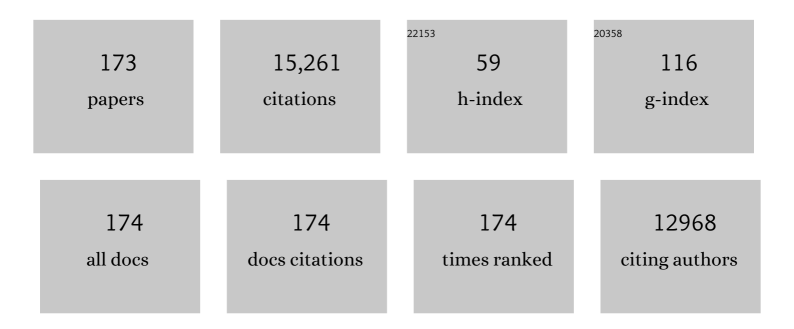
Shomi S Bhattacharya

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	<scp><i>WDR34</i></scp> , a candidate gene for nonâ€syndromic rodâ€cone dystrophy. Clinical Genetics, 2021, 99, 298-302.	2.0	7
2	Retinal pigment epithelium degeneration caused by aggregation of PRPF31 and the role of HSP70 family of proteins. Molecular Medicine, 2020, 26, 1.	4.4	45
3	Mutations in SPATA13/ASEF2 cause primary angle closure glaucoma. PLoS Genetics, 2020, 16, e1008721.	3.5	12
4	Subretinal Transplant of Induced Pluripotent Stem Cell-Derived Retinal Pigment Epithelium on Nanostructured Fibrin-Agarose. Tissue Engineering - Part A, 2019, 25, 799-808.	3.1	15
5	Generation and characterization of the human iPSC line CABi001-A from a patient with retinitis pigmentosa caused by a novel mutation in PRPF31 gene. Stem Cell Research, 2019, 36, 101426.	0.7	1
6	The Resveratrol Prodrug JC19 Delays Retinal Degeneration in rd10 Mice. Advances in Experimental Medicine and Biology, 2019, 1185, 457-462.	1.6	10
7	Rasagiline delays retinal degeneration in a mouse model of retinitis pigmentosa via modulation of Bax/Bclâ€2 expression. CNS Neuroscience and Therapeutics, 2018, 24, 448-455.	3.9	17
8	Identification and characterization of the VAX2 p.Leu139Arg variant: possible involvement of VAX2 in cone dystrophy. Ophthalmic Genetics, 2018, 39, 539-543.	1.2	1
9	Gene of the month: <i>PRPF31</i> . Journal of Clinical Pathology, 2017, 70, 729-732.	2.0	6
10	Effects of Ca2+ ions on bestrophin-1 surface films. Colloids and Surfaces B: Biointerfaces, 2017, 149, 226-232.	5.0	10
11	TOPORS, a Dual E3 Ubiquitin and Sumo1 Ligase, Interacts with 26 S Protease Regulatory Subunit 4, Encoded by the PSMC1 Gene. PLoS ONE, 2016, 11, e0148678.	2.5	10
12	EYS Is a Protein Associated with the Ciliary Axoneme in Rods and Cones. PLoS ONE, 2016, 11, e0166397.	2.5	36
13	Span poly-L-arginine nanoparticles are efficient non-viral vectors for PRPF31 gene delivery: An approach of gene therapy to treat retinitis pigmentosa. Nanomedicine: Nanotechnology, Biology, and Medicine, 2016, 12, 2251-2260.	3.3	18
14	Genome-wide association study identifies five new susceptibility loci for primary angle closure glaucoma. Nature Genetics, 2016, 48, 556-562.	21.4	147
15	A missense mutation inASRGL1is involved in causing autosomal recessive retinal degeneration. Human Molecular Genetics, 2016, 25, ddw113.	2.9	16
16	Transcriptional regulation of PRPF31 gene expression by MSR1 repeat elements causes incomplete penetrance in retinitis pigmentosa. Scientific Reports, 2016, 6, 19450.	3.3	42
17	First insights into the expression of VAX2 in humans and its localization in the adult primate retina. Experimental Eye Research, 2016, 148, 24-29.	2.6	7
18	267 Spanish Exomes Reveal Population-Specific Differences in Disease-Related Genetic Variation. Molecular Biology and Evolution, 2016, 33, 1205-1218.	8.9	78

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19	Genetic association and stress mediated down-regulation in trabecular meshwork implicates MPP7 as a novel candidate gene in primary open angle glaucoma. BMC Medical Genomics, 2016, 9, 15.	1.5	15
20	Human iPSC derived disease model of MERTK-associated retinitis pigmentosa. Scientific Reports, 2015, 5, 12910.	3.3	47
21	Genotype and Phenotype Studies in Autosomal Dominant Retinitis Pigmentosa (adRP) of the French Canadian Founder Population. , 2015, 56, 8297.		36
22	Cleavage of Mer Tyrosine Kinase (MerTK) from the Cell Surface Contributes to the Regulation of Retinal Phagocytosis. Journal of Biological Chemistry, 2015, 290, 4941-4952.	3.4	49
23	Concise Review: Reactive Astrocytes and Stem Cells in Spinal Cord Injury: Good Guys or Bad Guys?. Stem Cells, 2015, 33, 1036-1041.	3.2	108
24	Non-coding RNAs in pluripotency and neural differentiation of human pluripotent stem cells. Frontiers in Genetics, 2014, 5, 132.	2.3	22
25	ABCC5, a Gene That Influences the Anterior Chamber Depth, Is Associated with Primary Angle Closure Glaucoma. PLoS Genetics, 2014, 10, e1004089.	3.5	68
26	Dominant <i>PRPF31</i> Mutations Are Hypostatic to a Recessive <i>CNOT3</i> Polymorphism in Retinitis Pigmentosa: A Novel Phenomenon of "Linked <i>Trans</i> Acting Epistasis― Annals of Human Genetics, 2014, 78, 62-71.	0.8	28
27	The familial dementia gene revisited: a missense mutation revealed by whole-exome sequencing identifies ITM2B as a candidate gene underlying a novel autosomal dominant retinal dystrophy in a large family. Human Molecular Genetics, 2014, 23, 491-501.	2.9	29
28	Brief Report: Astrogliosis Promotes Functional Recovery of Completely Transected Spinal Cord Following Transplantation of hESC-Derived Oligodendrocyte and Motoneuron Progenitors. Stem Cells, 2014, 32, 594-599.	3.2	26
29	Mutations in Pre-mRNA Processing Factors 3, 8, and 31 Cause Dysfunction of the Retinal Pigment Epithelium. American Journal of Pathology, 2014, 184, 2641-2652.	3.8	62
30	Biallelic Variants in TTLL5, Encoding a Tubulin Glutamylase, Cause Retinal Dystrophy. American Journal of Human Genetics, 2014, 94, 760-769.	6.2	67
31	Rare and common variants in extracellular matrix gene Fibrillin 2 (FBN2) are associated with macular degeneration. Human Molecular Genetics, 2014, 23, 5827-5837.	2.9	52
32	Hypoxia Increases the Yield of Photoreceptors Differentiating from Mouse Embryonic Stem Cells and Improves the Modeling of Retinogenesis In Vitro. Stem Cells, 2013, 31, 966-978.	3.2	36
33	Wolfram gene (WFS1) mutation causes autosomal dominant congenital nuclear cataract in humans. European Journal of Human Genetics, 2013, 21, 1356-1360.	2.8	50
34	Whole-Exome Sequencing Identifies LRIT3 Mutations as a Cause of Autosomal-Recessive Complete Congenital Stationary Night Blindness. American Journal of Human Genetics, 2013, 92, 67-75.	6.2	120
35	A novel locus for autosomal dominant cone-rod dystrophy maps to chromosome 10q. European Journal of Human Genetics, 2013, 21, 338-342.	2.8	4
36	Novel <i>GUCA1A</i> Mutations Suggesting Possible Mechanisms of Pathogenesis in Cone, Cone-Rod, and Macular Dystrophy Patients. BioMed Research International, 2013, 2013, 1-15.	1.9	32

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37	Further Insights Into GPR179: Expression, Localization, and Associated Pathogenic Mechanisms Leading to Complete Congenital Stationary Night Blindness. , 2013, 54, 8041.		20
38	Disease-Causing Mutations in BEST1 Gene Are Associated with Altered Sorting of Bestrophin-1 Protein. International Journal of Molecular Sciences, 2013, 14, 15121-15140.	4.1	14
39	ATR localizes to the photoreceptor connecting cilium and deficiency leads to severe photoreceptor degeneration in mice. Human Molecular Genetics, 2013, 22, 1507-1515.	2.9	27
40	A Study into the Evolutionary Divergence of the Core Promoter Elements of PRPF31 and TFPT. Journal of Molecular and Genetic Medicine: an International Journal of Biomedical Research, 2013, 07, .	0.1	1
41	CNOT3 Is a Modifier of PRPF31 Mutations in Retinitis Pigmentosa with Incomplete Penetrance. PLoS Genetics, 2012, 8, e1003040.	3.5	109
42	Expression of PRPF31 and TFPT: regulation in health and retinal disease. Human Molecular Genetics, 2012, 21, 4126-4137.	2.9	15
43	Common Polymorphisms in theSERPINI2Gene Are Associated with Refractive Error in the 1958 British Birth Cohort. , 2012, 53, 440.		3
44	Hypoxia Enhances the Generation of Retinal Progenitor Cells from Human Induced Pluripotent and Embryonic Stem Cells. Stem Cells and Development, 2012, 21, 1344-1355.	2.1	51
45	Complement factor H genetic variant and age-related macular degeneration: effect size, modifiers and relationship to disease subtype. International Journal of Epidemiology, 2012, 41, 250-262.	1.9	79
46	NMNAT1 mutations cause Leber congenital amaurosis. Nature Genetics, 2012, 44, 1040-1045.	21.4	171
47	Cross species analysis of Prominin reveals a conserved cellular role in invertebrate and vertebrate photoreceptor cells. Developmental Biology, 2012, 371, 312-320.	2.0	41
48	Genome-wide association analyses identify three new susceptibility loci for primary angle closure glaucoma. Nature Genetics, 2012, 44, 1142-1146.	21.4	196
49	Development and application of a next-generation-sequencing (NGS) approach to detect known and novel gene defects underlying retinal diseases. Orphanet Journal of Rare Diseases, 2012, 7, 8.	2.7	144
50	Derivation of Cerebellar Neurons from Human Pluripotent Stem Cells. Current Protocols in Stem Cell Biology, 2012, 20, Unit 1H.5.	3.0	28
51	A map of human microRNA variation uncovers unexpectedly high levels of variability. Genome Medicine, 2012, 4, 62.	8.2	28
52	High Prevalence of Posterior Polymorphous Corneal Dystrophy in the Czech Republic; Linkage Disequilibrium Mapping and Dating an Ancestral Mutation. PLoS ONE, 2012, 7, e45495.	2.5	24
53	Whole-Exome Sequencing Identifies Mutations in GPR179 Leading to Autosomal-Recessive Complete Congenital Stationary Night Blindness. American Journal of Human Genetics, 2012, 90, 321-330.	6.2	121
54	RP1 and autosomal dominant rod-cone dystrophy: Novel mutations, a review of published variants, and genotype-phenotype correlation. Human Mutation, 2012, 33, 73-80.	2.5	33

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55	<i>CRB1</i> mutations in inherited retinal dystrophies. Human Mutation, 2012, 33, 306-315.	2.5	153
56	Three Gene-Targeted Mouse Models of RNA Splicing Factor RP Show Late-Onset RPE and Retinal Degeneration. , 2011, 52, 190.		70
57	A 112 kb Deletion in Chromosome 19q13.42 Leads to Retinitis Pigmentosa. , 2011, 52, 6597.		22
58	Concise Review: Stem Cells for the Treatment of Cerebellar-Related Disorders. Stem Cells, 2011, 29, 564-569.	3.2	7
59	Novel <i>C2orf71</i> mutations account for â^¼1% of cases in a large French arRP cohort. Human Mutation, 2011, 32, E2091-103.	2.5	29
60	TOPORS, implicated in retinal degeneration, is a cilia-centrosomal protein. Human Molecular Genetics, 2011, 20, 975-987.	2.9	49
61	Autosomal Dominant Retinitis Pigmentosa with Intrafamilial Variability and Incomplete Penetrance in Two Families Carrying Mutations in <i>PRPF8</i> ., 2011, 52, 9304.		38
62	Copy-Number Variations in <i>EYS:</i> A Significant Event in the Appearance of arRP., 2011, 52, 5625.		40
63	A novel 1-bp deletion in PITX3 causing congenital posterior polar cataract. Molecular Vision, 2011, 17, 1249-53.	1.1	16
64	Autosomal dominant Best disease with an unusual electrooculographic light rise and risk of angle-closure glaucoma: a clinical and molecular genetic study. Molecular Vision, 2011, 17, 2272-82.	1.1	16
65	RDH12 retinopathy: novel mutations and phenotypic description. Molecular Vision, 2011, 17, 2706-16.	1.1	47
66	Prognosis for splicing factor PRPF8 retinitis pigmentosa, novel mutations and correlation between human and yeast phenotypes. Human Mutation, 2010, 31, E1361-E1376.	2.5	31
67	EYS is a major gene for rod-cone dystrophies in France. Human Mutation, 2010, 31, E1406-E1435.	2.5	86
68	Mutation spectrum of EYS in Spanish patients with autosomal recessive retinitis pigmentosa. Human Mutation, 2010, 31, E1772-E1800.	2.5	69
69	Prevalence and novelty of PRPF31 mutations in French autosomal dominant rod-cone dystrophy patients and a review of published reports. BMC Medical Genetics, 2010, 11, 145.	2.1	49
70	Photoreceptor degeneration: genetic and mechanistic dissection of a complex trait. Nature Reviews Genetics, 2010, 11, 273-284.	16.3	519
71	Identification of Novel Mutations in the Ortholog of <i>Drosophila</i> Eyes Shut Gene (<i>EYS</i>) Causing Autosomal Recessive Retinitis Pigmentosa. , 2010, 51, 4266.		57
72	Spectrum of Rhodopsin Mutations in French Autosomal Dominant Rod–Cone Dystrophy Patients. , 2010, 51, 3687.		45

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73	Molecular genetic study of Egyptian patients with macular corneal dystrophy. British Journal of Ophthalmology, 2010, 94, 250-255.	3.9	10
74	Loss of lysophosphatidylcholine acyltransferase 1 leads to photoreceptor degeneration in <i>rd11</i> mice. Proceedings of the National Academy of Sciences of the United States of America, 2010, 107, 15523-15528.	7.1	55
75	Evidence for Keratoconus Susceptibility Locus on Chromosome 14. JAMA Ophthalmology, 2010, 128, 1191.	2.4	41
76	Novel mutations in MERTK associated with childhood onset rod-cone dystrophy. Molecular Vision, 2010, 16, 369-77.	1.1	73
77	Study of Gene-Targeted Mouse Models of Splicing Factor Gene <i>Prpf31</i> Implicated in Human Autosomal Dominant Retinitis Pigmentosa (RP). , 2009, 50, 5927.		52
78	Mutations in <i>TOPORS</i> : A Rare Cause of Autosomal Dominant Retinitis Pigmentosa in Continental Europe?. Ophthalmic Genetics, 2009, 30, 96-98.	1.2	8
79	Dominant cataract formation in association with a vimentin assembly disrupting mutation. Human Molecular Genetics, 2009, 18, 1052-1057.	2.9	88
80	Mutations of theEPHA2receptor tyrosine kinase gene cause autosomal dominant congenital cataract. Human Mutation, 2009, 30, E603-E611.	2.5	96
81	A common allele in RPGRIP1L is a modifier of retinal degeneration in ciliopathies. Nature Genetics, 2009, 41, 739-745.	21.4	255
82	Mutations in a BTB-Kelch Protein, KLHL7, Cause Autosomal-Dominant Retinitis Pigmentosa. American Journal of Human Genetics, 2009, 84, 792-800.	6.2	89
83	TRPM1 Is Mutated in Patients with Autosomal-Recessive Complete Congenital Stationary Night Blindness. American Journal of Human Genetics, 2009, 85, 720-729.	6.2	207
84	EYS, encoding an ortholog of Drosophila spacemaker, is mutated in autosomal recessive retinitis pigmentosa. Nature Genetics, 2008, 40, 1285-1287.	21.4	175
85	Effect of Gene Therapy on Visual Function in Leber's Congenital Amaurosis. New England Journal of Medicine, 2008, 358, 2231-2239.	27.0	1,793
86	Phenotype Associated with the H626P Mutation and Other Changes in the <i>TGFBI</i> Gene in Czech Families. Ophthalmic Research, 2008, 40, 105-108.	1.9	10
87	Dominant Cone and Cone-Rod Dystrophies: Functional Analysis of Mutations in RetGC1 and GCAP1. Novartis Foundation Symposium, 2008, 255, 37-50.	1.1	5
88	Disease mechanism for retinitis pigmentosa (RP11) caused by missense mutations in the splicing factor gene PRPF31. Molecular Vision, 2008, 14, 683-90.	1.1	26
89	Mutations in the Gene Coding for the Pre-mRNA Splicing Factor,PRPF31, in Patients with Autosomal Dominant Retinitis Pigmentosa. , 2007, 48, 1330.		60
90	An Assessment of the Apex Microarray Technology in Genotyping Patients with Leber Congenital Amaurosis and Early-Onset Severe Retinal Dystrophy. , 2007, 48, 5684.		56

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91	Mutations in splicing factor PRPF3, causing retinal degeneration, form detrimental aggregates in photoreceptor cells. Human Molecular Genetics, 2007, 16, 1699-1707.	2.9	46
92	A Clinical and Molecular Genetic Study of Egyptian and Saudi Arabian Patients With Primary Congenital Glaucoma (PCG). Journal of Glaucoma, 2007, 16, 104-111.	1.6	24
93	Mutations in TOPORS Cause Autosomal Dominant Retinitis Pigmentosa with Perivascular Retinal Pigment Epithelium Atrophy. American Journal of Human Genetics, 2007, 81, 1098-1103.	6.2	77
94	Novel mutations in theZEB1 gene identified in Czech and British patients with posterior polymorphous corneal dystrophy. Human Mutation, 2007, 28, 638-638.	2.5	67
95	The Roles of <i>PAX6</i> and <i>SOX2</i> in Myopia: Lessons from the 1958 British Birth Cohort. , 2007, 48, 4421.		37
96	Premature Truncation of a Novel Protein, RD3, Exhibiting Subnuclear Localization Is Associated with Retinal Degeneration. American Journal of Human Genetics, 2006, 79, 1059-1070.	6.2	112
97	A study of the nuclear trafficking of the splicing factor protein PRPF31 linked to autosomal dominant retinitis pigmentosa (ADRP). Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2006, 1762, 304-311.	3.8	18
98	Retinitis pigmentosa associated with rhodopsin mutations: Correlation between phenotypic variability and molecular effects. Vision Research, 2006, 46, 4556-4567.	1.4	76
99	Exclusion of Four Candidate Genes, <i>KHDRBS2, PTP4A1, KIAA1411</i> and <i>OGFRL1,</i> as Causative of Autosomal Recessive Retinitis Pigmentosa. Ophthalmic Research, 2006, 38, 19-23.	1.9	10
100	Maculopathy Due to the R345W Substitution in Fibulin-3: Distinct Clinical Features, Disease Variability, and Extent of Retinal Dysfunction. , 2006, 47, 3085.		48
101	A large deletion in the adRP gene PRPF31: evidence that haploinsufficiency is the cause of disease. Molecular Vision, 2006, 12, 384-8.	1.1	49
102	Molecular genetics of retinitis pigmentosa in two Romani (Gypsy) families. Molecular Vision, 2006, 12, 909-14.	1.1	20
103	A new locus (RP31) for autosomal dominant retinitis pigmentosa maps to chromosome 9p. Human Genetics, 2005, 118, 501-503.	3.8	13
104	Clinical Features and Course of Patients with Glaucoma with the E50K Mutation in the Optineurin Gene. , 2005, 46, 2816.		127
105	Posterior Polymorphous Corneal Dystrophy in Czech Families Maps to Chromosome 20 and Excludes theVSX1Gene. , 2005, 46, 4480.		67
106	Mutant carbonic anhydrase 4 impairs pH regulation and causes retinal photoreceptor degeneration. Human Molecular Genetics, 2005, 14, 255-265.	2.9	79
107	A Clinical and Molecular Genetic Study of Autosomal-Dominant Stromal Corneal Dystrophy in British Population. Ophthalmic Research, 2005, 37, 310-317.	1.9	16
108	Molecular Genetic Analysis of Two Functional Candidate Genes in the Autosomal Recessive Retinitis Pigmentosa, RP25, Locus. Current Eye Research, 2005, 30, 1081-1087.	1.5	8

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109	Novel CHST6 nonsense and missense mutations responsible for macular corneal dystrophy. American Journal of Ophthalmology, 2005, 139, 192-193.	3.3	28
110	Developmental Expression Profile of the Optic Atrophy Gene Product: OPA1 Is Not Localized Exclusively in the Mammalian Retinal Ganglion Cell Layer. , 2004, 45, 1667.		44
111	Mutations ofVMD2Splicing Regulators Cause Nanophthalmos and Autosomal Dominant Vitreoretinochoroidopathy (ADVIRC). , 2004, 45, 3683.		205
112	Purification, characterisation and intracellular localisation of aryl hydrocarbon interacting protein-like 1 (AIPL1) and effects of mutations associated with inherited retinal dystrophies. Biochimica Et Biophysica Acta - Molecular Basis of Disease, 2004, 1690, 141-149.	3.8	8
113	Molecular genetic basis of inherited cataract and associated phenotypes. Survey of Ophthalmology, 2004, 49, 300-315.	4.0	208
114	The Phenotype of Leber Congenital Amaurosis in Patients With AIPL1 Mutations. JAMA Ophthalmology, 2004, 122, 1029.	2.4	105
115	Fox's in development and disease. Trends in Genetics, 2003, 19, 339-344.	6.7	316
116	Expression ofPRPF31mRNA in Patients with Autosomal Dominant Retinitis Pigmentosa: A Molecular Clue for Incomplete Penetrance?. , 2003, 44, 4204.		125
117	Phenotype of Retinitis Pigmentosa Associated With the Ser50Thr Mutation in the NRL Gene. JAMA Ophthalmology, 2003, 121, 793.	2.4	21
118	Disease mechanism for retinitis pigmentosa (RP11) caused by mutations in the splicing factor gene PRPF31. Human Molecular Genetics, 2002, 11, 3209-3219.	2.9	75
119	Mutations in HPRP3, a third member ofpre-mRNA splicing factor genes, implicated in autosomal dominant retinitis pigmentosa. Human Molecular Genetics, 2002, 11, 87-92.	2.9	217
120	Mutations in a protein target of the Pim-1 kinase associated with the RP9 form of autosomal dominant retinitis pigmentosa. European Journal of Human Genetics, 2002, 10, 245-249.	2.8	87
121	The ABCA4 2588G>C Stargardt mutation: single origin and increasing frequency from South-West to North-East Europe. European Journal of Human Genetics, 2002, 10, 197-203.	2.8	45
122	An Integrated, Functionally Annotated Gene Map of the DXS8026–ELK1 Interval on Human Xp11.3–Xp11.23: Potential Hotspot for Neurogenetic Disorders. Genomics, 2002, 79, 560-572.	2.9	103
123	Reply to Veromann. American Journal of Human Genetics, 2002, 71, 685-686.	6.2	0
124	Characterisation of two genes for guanylate cyclase activator protein (GCAP1 and GCAP2) in the Japanese pufferfish, Fugu rubripes. Biochimica Et Biophysica Acta Gene Regulatory Mechanisms, 2002, 1577, 73-80.	2.4	1
125	Cloning and characterization of WDR17, a novel WD repeat-containing gene on chromosome 4q34. Biochimica Et Biophysica Acta Gene Regulatory Mechanisms, 2002, 1579, 18-25.	2.4	12
126	A major marker for normal tension glaucoma: association with polymorphisms in the OPA1 gene. Human Genetics, 2002, 110, 52-56.	3.8	123

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127	Investigating the association between OPA1 polymorphisms and glaucoma: comparison between normal tension and high tension primary open angle glaucoma. Human Genetics, 2002, 110, 513-514.	3.8	49
128	Identification of novel mutations in the carbohydrate sulfotransferase gene (CHST6) causing macular corneal dystrophy. Investigative Ophthalmology and Visual Science, 2002, 43, 377-82.	3.3	36
129	Ocular developmental abnormalities and glaucoma associated with interstitial 6p25 duplications and deletions. Investigative Ophthalmology and Visual Science, 2002, 43, 1843-9.	3.3	63
130	Identification and Functional Consequences of a New Mutation (E155G) in the Gene for GCAP1 That Causes Autosomal Dominant Cone Dystrophy. American Journal of Human Genetics, 2001, 69, 471-480.	6.2	115
131	Alpha-B Crystallin Gene (CRYAB) Mutation Causes Dominant Congenital Posterior Polar Cataract in Humans. American Journal of Human Genetics, 2001, 69, 1141-1145.	6.2	208
132	Molecular genetics and prospects for therapy of the inherited retinal dystrophies. Current Opinion in Genetics and Development, 2001, 11, 307-316.	3.3	92
133	A Human Homolog of Yeast Pre-mRNA Splicing Gene, PRP31, Underlies Autosomal Dominant Retinitis Pigmentosa on Chromosome 19q13.4 (RP11). Molecular Cell, 2001, 8, 375-381.	9.7	305
134	A frameshift mutation in exon 28 of the OPA1 gene explains the high prevalence of dominant optic atrophy in the Danish population: evidence for a founder effect. Human Genetics, 2001, 109, 498-502.	3.8	59
135	Novel mutations of theRPGR gene in RP3 families. Human Mutation, 2000, 15, 386-386.	2.5	12
136	Novel frameshift mutations in theRP2 gene and polymorphic variants. Human Mutation, 2000, 15, 580-580.	2.5	22
137	Sequence variation within theRPGR gene: Evidence for a founder complex allele. Human Mutation, 2000, 16, 273-274.	2.5	8
138	Mutations in a new photoreceptor-pineal gene on 17p cause Leber congenital amaurosis. Nature Genetics, 2000, 24, 79-83.	21.4	257
139	Restoration of photoreceptor ultrastructure and function in retinal degeneration slow mice by gene therapy. Nature Genetics, 2000, 25, 306-310.	21.4	295
140	OPA1, encoding a dynamin-related GTPase, is mutated in autosomal dominant optic atrophy linked to chromosome 3q28. Nature Genetics, 2000, 26, 211-215.	21.4	1,169
141	NRL S50T mutation and the importance of â€~founder effects' in inherited retinal dystrophies. European Journal of Human Genetics, 2000, 8, 783-787.	2.8	18
142	Functional characterization of missense mutations at codon 838 in retinal guanylate cyclase correlates with disease severity in patients with autosomal dominant cone-rod dystrophy. Human Molecular Genetics, 2000, 9, 3065-3073.	2.9	83
143	Characterization of the Human TBX20 Gene, a New Member of the T-Box Gene Family Closely Related to the Drosophila H15 Gene. Genomics, 2000, 67, 317-332.	2.9	44
144	Prevalence of AIPL1 Mutations in Inherited Retinal Degenerative Disease. Molecular Genetics and Metabolism, 2000, 70, 142-150.	1.1	144

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145	Chromosomal Duplication Involving the Forkhead Transcription Factor Gene FOXC1 Causes Iris Hypoplasia and Glaucoma. American Journal of Human Genetics, 2000, 67, 1129-1135.	6.2	105
146	Chromosomal Duplication Involving the Forkhead Transcription Factor Gene <i>FOXC1</i> Causes Iris Hypoplasia and Glaucoma. American Journal of Human Genetics, 2000, 67, 1129-1135.	6.2	127
147	Mutations in a human homologue of Drosophila crumbs cause retinitis pigmentosa (RP12). Nature Genetics, 1999, 23, 217-221.	21.4	427
148	A mutation in NRL is associated with autosomal dominant retinitis pigmentosa. Nature Genetics, 1999, 21, 355-356.	21.4	205
149	Connexin 50 mutation in a family with congenital "zonular nuclear" pulverulent cataract of Pakistani origin. Human Genetics, 1999, 105, 168-170.	3.8	33
150	Connexin 50 mutation in a family with congenital "zonular nuclear" pulverulent cataract of Pakistani origin. Human Genetics, 1999, 105, 168-170.	3.8	139
151	Refinement of the Locus for Autosomal Recessive Retinitis Pigmentosa (RP25) Linked to Chromosome 6q in a Family of Pakistani Origin. American Journal of Human Genetics, 1999, 65, 571-574.	6.2	32
152	Characterization of the human diacylglycerol kinase ε gene and its assessment as a candidate for inherited retinitis pigmentosa. Gene, 1999, 239, 185-192.	2.2	7
153	Dominant optic atrophy: exclusion and fine genetic mapping of the candidate gene, HRY. Mammalian Genome, 1998, 9, 784-787.	2.2	5
154	High frequency of persistent hyperplastic primary vitreous and cataracts in p53-deficient mice. Cell Death and Differentiation, 1998, 5, 156-162.	11.2	67
155	Genetic blindness: current concepts in the pathogenesis of human outer retinal dystrophies. Trends in Genetics, 1998, 14, 103-108.	6.7	60
156	Demonstration of a founder effect and fine mapping of dominant optic atrophy locus on 3q28-qter by linkage disequilibrium method. Human Genetics, 1998, 102, 79-86.	3.8	31
157	GCAP1(Y99C) Mutant Is Constitutively Active in Autosomal Dominant Cone Dystrophy. Molecular Cell, 1998, 2, 129-133.	9.7	150
158	Founder Effect, Seen in the British Population, of the 172 Peripherin/RDS Mutation—and Further Refinement of Genetic Positioning of the Peripherin/RDS Gene. American Journal of Human Genetics, 1998, 62, 192-195.	6.2	42
159	Segregation of a PRKCG Mutation in Two RP11 Families. American Journal of Human Genetics, 1998, 62, 1248-1252.	6.2	21
160	A Locus for Autosomal Recessive Congenital Microphthalmia Maps to Chromosome 14q32. American Journal of Human Genetics, 1998, 62, 1113-1116.	6.2	41
161	Absence of p53 delays apoptotic photoreceptor cell death in the rds mouse. Current Eye Research, 1998, 17, 917-923.	1.5	23
162	Adeno-Associated Virus Gene Transfer to Mouse Retina. Human Gene Therapy, 1998, 9, 81-86.	2.7	118

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#	Article	IF	CITATIONS
163	Cone-Rod Dystrophy Due to Mutations in a Novel Photoreceptor-Specific Homeobox Gene (CRX) Essential for Maintenance of the Photoreceptor. Cell, 1997, 91, 543-553.	28.9	520
164	Gene transfer into the mouse retina mediated by an adeno-associated viral vector. Human Molecular Genetics, 1996, 5, 591-594.	2.9	209
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