Michael E Cheetham

List of Publications by Year in descending order

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156 papers 18,210 citations

28274 55 h-index 128 g-index

164 all docs

164 docs citations

164 times ranked 26094 citing authors

#	Article	IF	CITATIONS
1	Induced Pluripotent Stem Cells for Inherited Optic Neuropathies—Disease Modeling and Therapeutic Development. Journal of Neuro-Ophthalmology, 2022, 42, 35-44.	0.8	12
2	Eye on a Dish Models to Evaluate Splicing Modulation. Methods in Molecular Biology, 2022, 2434, 245-255.	0.9	5
3	Intravitreal antisense oligonucleotide sepofarsen in Leber congenital amaurosis type 10: a phase 1b/2 trial. Nature Medicine, 2022, 28, 1014-1021.	30.7	46
4	Modelling autosomal dominant optic atrophy associated with <i>OPA1</i> variants in iPSC-derived retinal ganglion cells. Human Molecular Genetics, 2022, 31, 3478-3493.	2.9	9
5	Allele-specific editing ameliorates dominant retinitis pigmentosa in a transgenic mouse model. American Journal of Human Genetics, 2021, 108, 295-308.	6.2	31
6	A look into retinal organoids: methods, analytical techniques, and applications. Cellular and Molecular Life Sciences, 2021, 78, 6505-6532.	5.4	36
7	CRISPR-Cas9 correction of OPA1 c.1334G>A: p.R445H restores mitochondrial homeostasis in dominant optic atrophy patient-derived iPSCs. Molecular Therapy - Nucleic Acids, 2021, 26, 432-443.	5.1	21
8	Guidelines for the use and interpretation of assays for monitoring autophagy (4th) Tj ETQq0 0 0 rgBT /Overlock	10 Jf 50 4	62 Td (edition
9	Negative Regulator of Ubiquitin-Like Protein 1 modulates the autophagy–lysosomal pathway via p62 to facilitate the extracellular release of tau following proteasome impairment. Human Molecular Genetics, 2020, 29, 80-96.	2.9	12
10	Detailed Phenotyping and Therapeutic Strategies for Intronic ABCA4 Variants in Stargardt Disease. Molecular Therapy - Nucleic Acids, 2020, 21, 412-427.	5.1	55
11	Structural Variants Create New Topological-Associated Domains and Ectopic Retinal Enhancer-Gene Contact in Dominant Retinitis Pigmentosa. American Journal of Human Genetics, 2020, 107, 802-814.	6.2	75
12	DYNC2H1 hypomorphic or retina-predominant variants cause nonsyndromic retinal degeneration. Genetics in Medicine, 2020, 22, 2041-2051.	2.4	38
13	Modeling and Rescue of RP2 Retinitis Pigmentosa Using iPSC-Derived Retinal Organoids. Stem Cell Reports, 2020, 15, 67-79.	4.8	109
14	Symmetric arrangement of mitochondria:plasma membrane contacts between adjacent photoreceptor cells regulated by Opa1. Proceedings of the National Academy of Sciences of the United States of America, 2020, 117, 15684-15693.	7.1	26
15	AAV-mediated ERdj5 overexpression protects against P23H rhodopsin toxicity. Human Molecular Genetics, 2020, 29, 1310-1318.	2.9	10
16	PCARE and WASF3 regulate ciliary F-actin assembly that is required for the initiation of photoreceptor outer segment disk formation. Proceedings of the National Academy of Sciences of the United States of America, 2020, 117, 9922-9931.	7.1	58
17	Delineating the expanding phenotype associated with <i>SCAPER</i> gene mutation. American Journal of Medical Genetics, Part A, 2019, 179, 1665-1671.	1.2	10
18	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.	5.2	25

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19	Effect of an intravitreal antisense oligonucleotide on vision in Leber congenital amaurosis due to a photoreceptor cilium defect. Nature Medicine, 2019, 25, 225-228.	30.7	177
20	Antisense Therapy for a Common Corneal Dystrophy Ameliorates TCF4 Repeat Expansion-Mediated Toxicity. American Journal of Human Genetics, 2018, 102, 528-539.	6.2	59
21	Ectopic GRHL2 Expression Due to Non-coding Mutations Promotes Cell State Transition and Causes Posterior Polymorphous Corneal Dystrophy 4. American Journal of Human Genetics, 2018, 102, 447-459.	6.2	45
22	A recurrent splice-site mutation in <i>EPHA2</i> causing congenital posterior nuclear cataract. Ophthalmic Genetics, 2018, 39, 236-241.	1.2	13
23	Missense variants in the X-linked gene <i>PRPS1</i> cause retinal degeneration in females. Human Mutation, 2018, 39, 80-91.	2.5	23
24	The molecular and cellular basis of rhodopsin retinitis pigmentosa reveals potential strategies for therapy. Progress in Retinal and Eye Research, 2018, 62, 1-23.	15.5	254
25	Whole-genome sequencing reveals a recurrent missense mutation in the Connexin 46 (GJA3) gene causing autosomal-dominant lamellar cataract. Eye, 2018, 32, 1661-1668.	2.1	6
26	Splice-Modulating Oligonucleotide QR-110 Restores CEP290 mRNA and Function in Human c.2991+1655A>G LCA10 Models. Molecular Therapy - Nucleic Acids, 2018, 12, 730-740.	5.1	130
27	DNAJ Proteins in neurodegeneration: essential and protective factors. Philosophical Transactions of the Royal Society B: Biological Sciences, 2018, 373, 20160534.	4.0	115
28	Correlative light and immuno-electron microscopy of retinal tissue cryostat sections. PLoS ONE, 2018, 13, e0191048.	2.5	12
29	Novel homozygous splicing mutations in cause autosomal recessive retinitis pigmentosa. Molecular Vision, 2018, 24, 603-612.	1.1	6
30	Protein kinase CK2 modulates HSJ1 function through phosphorylation of the UIM2 domain. Human Molecular Genetics, 2017, 26, ddw420.	2.9	8
31	Biallelic Mutation of ARHGEF18, Involved in the Determination of Epithelial Apicobasal Polarity, Causes Adult-Onset Retinal Degeneration. American Journal of Human Genetics, 2017, 100, 334-342.	6.2	26
32	Rescue of mutant rhodopsin traffic by metformin-induced AMPK activation accelerates photoreceptor degeneration. Human Molecular Genetics, 2017, 26, ddw387.	2.9	39
33	Mutations in the Spliceosome Component CWC27 Cause Retinal Degeneration with or without Additional Developmental Anomalies. American Journal of Human Genetics, 2017, 100, 592-604.	6.2	61
34	REEP6 deficiency leads to retinal degeneration through disruption of ER homeostasis and protein trafficking. Human Molecular Genetics, 2017, 26, 2667-2677.	2.9	39
35	Arl3 and RP2 regulate the trafficking of ciliary tip kinesins. Human Molecular Genetics, 2017, 26, 2480-2492.	2.9	60
36	Pheno4J: a gene to phenotype graph database. Bioinformatics, 2017, 33, 3317-3319.	4.1	9

3

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37	Rescue of the MERTK phagocytic defect in a human iPSC disease model using translational read-through inducing drugs. Scientific Reports, 2017, 7, 51.	3.3	55
38	The role of the ER stress-response protein PERK in rhodopsin retinitis pigmentosa. Human Molecular Genetics, 2017, 26, 4896-4905.	2.9	82
39	Using induced pluripotent stem cells to understand retinal ciliopathy disease mechanisms and develop therapies. Biochemical Society Transactions, 2016, 44, 1245-1251.	3.4	18
40	Identification and Correction of Mechanisms Underlying Inherited Blindness in Human iPSC-Derived Optic Cups. Cell Stem Cell, 2016, 18, 769-781.	11.1	279
41	Recessive Retinopathy Consequent on Mutant G-Protein β Subunit 3 (<i>GNB3</i>). JAMA Ophthalmology, 2016, 134, 924.	2.5	25
42	Mutations in REEP6 Cause Autosomal-Recessive Retinitis Pigmentosa. American Journal of Human Genetics, 2016, 99, 1305-1315.	6.2	121
43	Autosomal-Dominant Corneal Endothelial Dystrophies CHED1 and PPCD1 Are Allelic Disorders Caused by Non-coding Mutations in the Promoter of OVOL2. American Journal of Human Genetics, 2016, 98, 75-89.	6.2	70
44	Guidelines for the use and interpretation of assays for monitoring autophagy (3rd edition). Autophagy, 2016, 12, 1-222.	9.1	4,701
45	The heat shock response plays an important role in TDP-43 clearance: evidence for dysfunction in amyotrophic lateral sclerosis. Brain, 2016, 139, 1417-1432.	7.6	131
46	Hsp90 as a Potential Therapeutic Target in Retinal Disease. Advances in Experimental Medicine and Biology, 2016, 854, 161-167.	1.6	19
47	Targeting the Proteostasis Network in Rhodopsin Retinitis Pigmentosa. Advances in Experimental Medicine and Biology, 2016, 854, 479-484.	1.6	7
48	Lack of Interphotoreceptor Retinoid Binding Protein Caused by Homozygous Mutation of <i>RBP3 </i> Is Associated With High Myopia and Retinal Dystrophy., 2015, 56, 2358.		42
49	Brittle Cornea Syndrome ZNF469 Mutation Carrier Phenotype and Segregation Analysis of Rare ZNF469 Variants in Familial Keratoconus. Investigative Ophthalmology and Visual Science, 2015, 56, 578-586.	3.3	33
50	Translational read-through of the RP2 Arg120stop mutation in patient iPSC-derived retinal pigment epithelium cells. Human Molecular Genetics, 2015, 24, 972-986.	2.9	97
51	Biallelic Mutations in the Autophagy Regulator DRAM2 Cause Retinal Dystrophy with Early Macular Involvement. American Journal of Human Genetics, 2015, 96, 948-954.	6.2	42
52	Molecular chaperones and neuronal proteostasis. Seminars in Cell and Developmental Biology, 2015, 40, 142-152.	5.0	81
53	The Role of HSP70 and Its Co-chaperones in Protein Misfolding, Aggregation and Disease. Sub-Cellular Biochemistry, 2015, 78, 243-273.	2.4	76
54	The Leber Congenital Amaurosis Protein AIPL1 and EB Proteins Co-Localize at the Photoreceptor Cilium. PLoS ONE, 2015, 10, e0121440.	2.5	8

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55	Differential Light-induced Responses in Sectorial Inherited Retinal Degeneration. Journal of Biological Chemistry, 2014, 289, 35918-35928.	3.4	32
56	The heat-shock response co-inducer arimoclomol protects against retinal degeneration in rhodopsin retinitis pigmentosa. Cell Death and Disease, 2014, 5, e1236-e1236.	6.3	54
57	The co-chaperone and reductase ERdj5 facilitates rod opsin biogenesis and quality control. Human Molecular Genetics, 2014, 23, 6594-6606.	2.9	23
58	Hsp90 inhibition protects against inherited retinal degeneration. Human Molecular Genetics, 2014, 23, 2164-2175.	2.9	70
59	Three Different Cone Opsin Gene Array Mutational Mechanisms; Genotype-Phenotype Correlation and Functional Investigation of Cone Opsin Variants. Human Mutation, 2014, 35, n/a-n/a.	2.5	53
60	Mutations in ARL2BP, Encoding ADP-Ribosylation-Factor-Like 2 Binding Protein, Cause Autosomal-Recessive Retinitis Pigmentosa. American Journal of Human Genetics, 2013, 93, 321-329.	6.2	67
61	The cell stress machinery and retinal degeneration. FEBS Letters, 2013, 587, 2008-2017.	2.8	126
62	Molecular Chaperone Mediated Late-Stage Neuroprotection in the SOD1G93A Mouse Model of Amyotrophic Lateral Sclerosis. PLoS ONE, 2013, 8, e73944.	2.5	51
63	Unpicking the UPR. , 2012, 53, 7167.		O
64	Deep intronic mutation in OFD1, identified by targeted genomic next-generation sequencing, causes a severe form of X-linked retinitis pigmentosa (RP23). Human Molecular Genetics, 2012, 21, 3647-3654.	2.9	133
65	The X-linked retinitis pigmentosa protein RP2 facilitates G protein traffic. Human Molecular Genetics, 2012, 21, 863-873.	2.9	34
66	Suppression of protein aggregation by chaperone modification of high molecular weight complexes. Brain, 2012, 135, 1180-1196.	7.6	103
67	NUB1 modulation of GSK3β reduces tau aggregation. Human Molecular Genetics, 2012, 21, 5254-5267.	2.9	29
68	BiP prevents rod opsin aggregation. Molecular Biology of the Cell, 2012, 23, 3522-3531.	2.1	44
69	Arl3 and RP2 mediated assembly and traffic of membrane associated cilia proteins. Vision Research, 2012, 75, 2-4.	1.4	51
70	The Inherited Blindness Protein AIPL1 Regulates the Ubiquitin-Like FAT10 Pathway. PLoS ONE, 2012, 7, e30866.	2.5	17
71	X-Linked Megalocornea Caused by Mutations in CHRDL1 Identifies an Essential Role for Ventroptin in Anterior Segment Development. American Journal of Human Genetics, 2012, 90, 247-259.	6.2	59
72	The Role of the X-linked Retinitis Pigmentosa Protein RP2 in Vesicle Traffic and Cilia Function. Advances in Experimental Medicine and Biology, 2012, 723, 527-532.	1.6	6

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73	A Novel Missense Mutation in Both OPN1LW and OPN1MW Cone Opsin Genes Causes X-Linked Cone Dystrophy (XLCOD5). Advances in Experimental Medicine and Biology, 2012, 723, 595-601.	1.6	2
74	Leber Congenital Amaurosis Associated with AIPL1: Challenges in Ascribing Disease Causation, Clinical Findings, and Implications for Gene Therapy. PLoS ONE, 2012, 7, e32330.	2.5	28
75	Functional characterization of a novel c.614-622del rhodopsin mutation in a French pedigree with retinitis pigmentosa. Molecular Vision, 2012, 18, 581-7.	1.1	2
76	Focus on Molecules: Centrosomal protein 290 (CEP290). Experimental Eye Research, 2011, 92, 316-317.	2.6	15
77	A simple cell based assay to measure Parkin activity. Journal of Neurochemistry, 2011, 116, 342-349.	3.9	5
78	Knockdown of the Zebrafish Ortholog of the Retinitis Pigmentosa 2 (<i>RP2</i>) Gene Results in Retinal Degeneration., 2011, 52, 2960.		32
79	Molecular chaperone-mediated rescue of mitophagy by a Parkin RING1 domain mutant. Human Molecular Genetics, 2011, 20, 16-27.	2.9	63
80	Molecular Mechanisms of Disease for Mutations at Gly-90 in Rhodopsin. Journal of Biological Chemistry, 2011, 286, 39993-40001.	3.4	37
81	TOPORS, implicated in retinal degeneration, is a cilia-centrosomal protein. Human Molecular Genetics, 2011, 20, 975-987.	2.9	49
82	X-Linked Cone Dystrophy Caused by Mutation of the Red and Green Cone Opsins. American Journal of Human Genetics, 2010, 87, 26-39.	6.2	45
83	COMMENTARY: Chaperoning against neuronal vulnerability (Commentary on Zijlstra <i>et al.</i>). European Journal of Neuroscience, 2010, 32, 759-759.	2.6	0
84	The Nanceâ€"Horan syndrome protein encodes a functional WAVE homology domain (WHD) and is important for co-ordinating actin remodelling and maintaining cell morphology. Human Molecular Genetics, 2010, 19, 2421-2432.	2.9	64
85	The retinitis pigmentosa protein RP2 links pericentriolar vesicle transport between the Golgi and the primary cilium. Human Molecular Genetics, 2010, 19, 1358-1367.	2.9	124
86	The binding of the molecular chaperone Hsc70 to the prion protein PrP is modulated by pH and copper. International Journal of Biochemistry and Cell Biology, 2010, 42, 1226-1232.	2.8	8
87	Pharmacological Manipulation of Rhodopsin Retinitis Pigmentosa. Advances in Experimental Medicine and Biology, 2010, 664, 317-323.	1.6	23
88	A dual role for EDEM1 in the processing of rod opsin. Journal of Cell Science, 2009, 122, 4465-4472.	2.0	52
89	Differential Expression of Two Distinct Functional Isoforms of Melanopsin (<i>Opn4</i>) in the Mammalian Retina. Journal of Neuroscience, 2009, 29, 12332-12342.	3.6	87
90	The ataxia protein sacsin is a functional co-chaperone that protects against polyglutamine-expanded ataxin-1. Human Molecular Genetics, 2009, 18, 1556-1565.	2.9	153

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91	DnaJB6 is present in the core of Lewy bodies and is highly upâ€regulated in parkinsonian astrocytes. Journal of Neuroscience Research, 2009, 87, 238-245.	2.9	67
92	Guidelines for the nomenclature of the human heat shock proteins. Cell Stress and Chaperones, 2009, 14, 105-111.	2.9	1,105
93	Distinct Melanocortin 2 Receptor Accessory Protein Domains Are Required for Melanocortin 2 Receptor Interaction and Promotion of Receptor Trafficking. Endocrinology, 2009, 150, 720-726.	2.8	63
94	MRAP and MRAP2 are bidirectional regulators of the melanocortin receptor family. Proceedings of the National Academy of Sciences of the United States of America, 2009, 106, 6146-6151.	7.1	201
95	EYS, encoding an ortholog of Drosophila spacemaker, is mutated in autosomal recessive retinitis pigmentosa. Nature Genetics, 2008, 40, 1285-1287.	21.4	175
96	Late stage treatment with arimoclomol delays disease progression and prevents protein aggregation in the SOD1 ^{G93A} mouse model of ALS. Journal of Neurochemistry, 2008, 107, 339-350.	3.9	147
97	Molecular chaperones and photoreceptor function. Progress in Retinal and Eye Research, 2008, 27, 434-449.	15.5	52
98	Pharmacological manipulation of gain-of-function and dominant-negative mechanisms in rhodopsin retinitis pigmentosa. Human Molecular Genetics, 2008, 17, 3043-3054.	2.9	125
99	The Melanocortin 2 Receptor Accessory Protein Exists as a Homodimer and Is Essential for the Function of the Melanocortin 2 Receptor in the Mouse Y1 Cell Line. Endocrinology, 2008, 149, 1935-1941.	2.8	65
100	The Leber Congenital Amaurosis Protein AIPL1 Functions as Part of a Chaperone Heterocomplex. , 2008, 49, 2878.		63
101	The Role of Chaperones and Co-Chaperones in Retinal Degenerative Diseases. , 2008, , 109-123.		0
102	Calnexin is not essential for mammalian rod opsin biogenesis. Molecular Vision, 2008, 14, 2466-74.	1.1	21
103	Hsp40 Molecules That Target to the Ubiquitin-proteasome System Decrease Inclusion Formation in Models of Polyglutamine Disease. Molecular Therapy, 2007, 15, 1100-1105.	8.2	96
104	An Assessment of the Apex Microarray Technology in Genotyping Patients with Leber Congenital Amaurosis and Early-Onset Severe Retinal Dystrophy., 2007, 48, 5684.		56
105	Targeting amyloid- \hat{l}^2 in glaucoma treatment. Proceedings of the National Academy of Sciences of the United States of America, 2007, 104, 13444-13449.	7.1	315
106	Cytosolic and ER J-domains of mammalian and parasitic origin can functionally interact with DnaK. International Journal of Biochemistry and Cell Biology, 2007, 39, 736-751.	2.8	32
107	Novel Retinal and Cone Photoreceptor Transcripts Revealed by Human Macular Expression Profiling. , 2007, 48, 5388.		26
108	Mutations in LCA5, encoding the ciliary protein lebercilin, cause Leber congenital amaurosis. Nature Genetics, 2007, 39, 889-895.	21.4	186

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109	The Role of Hsp70 and Its Co-Chaperones in Protein Misfolding, Aggregation and Disease. , 2007, , 122-136.		0
110	The retinitis pigmentosa-mutated RP2 protein exhibits exonuclease activity and translocates to the nucleus in response to DNA damage. Experimental Cell Research, 2006, 312, 1323-1334.	2.6	23
111	Focus on Molecules: Xâ€linked Retinitis Pigmentosa 2 protein, RP2. Experimental Eye Research, 2006, 82, 543-544.	2.6	20
112	The Chaperone Function of the LCA Protein AIPL1. Advances in Experimental Medicine and Biology, 2006, 572, 471-476.	1.6	4
113	Cystamine and cysteamine increase brain levels of BDNF in Huntington disease via HSJ1b and transglutaminase. Journal of Clinical Investigation, 2006, 116, 1410-1424.	8.2	211
114	Not all J domains are created equal: Implications for the specificity of Hsp40-Hsp70 interactions. Protein Science, 2005, 14, 1697-1709.	7.6	265
115	Treatment with extracellular HSP70/HSC70 protein can reduce polyglutamine toxicity and aggregation. Journal of Neurochemistry, 2005, 94, 597-606.	3.9	74
116	Mutations in MRAP, encoding a new interacting partner of the ACTH receptor, cause familial glucocorticoid deficiency type 2. Nature Genetics, 2005, 37, 166-170.	21.4	388
117	HSJ1 Is a Neuronal Shuttling Factor for the Sorting of Chaperone Clients to the Proteasome. Current Biology, 2005, 15, 1058-1064.	3.9	130
118	Identification of NovelRPGRORF15 Mutations in X-linked Progressive Cone-Rod Dystrophy (XLCORD) Families. , 2005, 46, 1891.		104
119	Assay and Functional Analysis of the ARL3 Effector RP2 Involved in Xâ€Linked Retinitis Pigmentosa. Methods in Enzymology, 2005, 404, 468-480.	1.0	6
120	Mechanisms of cell death in rhodopsin retinitis pigmentosa: implications for therapy. Trends in Molecular Medicine, 2005, 11, 177-185.	6.7	339
121	Isolation and characterization of murine Cds (CDP-diacylglycerol synthase) 1 and 2. Gene, 2005, 356, 19-31.	2.2	45
122	Focus on Molecules: Nyctalopin. Experimental Eye Research, 2005, 81, 627-628.	2.6	9
123	Inherited ACTH insensitivity illuminates the mechanisms of ACTH action. Trends in Endocrinology and Metabolism, 2005, 16, 451-457.	7.1	7 5
124	Novel CHST6 nonsense and missense mutations responsible for macular corneal dystrophy. American Journal of Ophthalmology, 2005, 139, 192-193.	3.3	28
125	Predominant rod photoreceptor degeneration in Leber congenital amaurosis. Molecular Vision, 2005, 11, 542-53.	1.1	12
126	Downstream caspases are novel targets for the antiapoptotic activity of the molecular chaperone Hsp70. Cell Stress and Chaperones, 2004, 9, 265.	2.9	70

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127	Nuclear translocation of the Hsp70/Hsp90 organizing protein mSTI1 is regulated by cell cycle kinases. Journal of Cell Science, 2004, 117, 701-710.	2.0	100
128	Modulation of Sub-RPE Deposits In Vitro: A Potential Model for Age-Related Macular Degeneration. Investigative Ophthalmology and Visual Science, 2004, 45, 1281-1288.	3.3	22
129	The Leber Congenital Amaurosis Protein AIPL1 Modulates the Nuclear Translocation of NUB1 and Suppresses Inclusion Formation by NUB1 Fragments. Journal of Biological Chemistry, 2004, 279, 48038-48047.	3.4	30
130	Oxidative Stress Affects the Junctional Integrity of Retinal Pigment Epithelial Cells., 2004, 45, 675.		229
131	Neuronal DnaJ proteins HSJ1a and HSJ1b: a role in linking the Hsp70 chaperone machine to the ubiquitin–proteasome system?. Biochemical Society Transactions, 2004, 32, 640-642.	3.4	36
132	Mutations in the CACNA1F and NYX genes in British CSNBX families. Human Mutation, 2003, 21, 169-169.	2.5	48
133	The Expression of the Leber Congenital Amaurosis Protein AIPL1 Coincides with Rod and Cone Photoreceptor Development. , 2003, 44, 5396.		45
134	The Chaperone Environment at the Cytoplasmic Face of the Endoplasmic Reticulum Can Modulate Rhodopsin Processing and Inclusion Formation. Journal of Biological Chemistry, 2003, 278, 19087-19094.	3.4	99
135	Organization on the plasma membrane of the retinitis pigmentosa protein RP2: investigation of association with detergent-resistant membranes and polarized sorting. Biochemical Journal, 2003, 372, 427-433.	3.7	23
136	Localization in the human retina of the X-linked retinitis pigmentosa protein RP2, its homologue cofactor C and the RP2 interacting protein Arl3. Human Molecular Genetics, 2002, 11, 3065-3074.	2.9	119
137	The Leber congenital amaurosis gene product AIPL1 is localized exclusively in rod photoreceptors of the adult human retina. Human Molecular Genetics, 2002, 11, 823-831.	2.9	67
138	The cellular fate of mutant rhodopsin: quality control, degradation and aggresome formation. Journal of Cell Science, 2002, 115, 2907-2918.	2.0	295
139	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
140	Delineation of the plasma membrane targeting domain of the X-linked retinitis pigmentosa protein RP2. Investigative Ophthalmology and Visual Science, 2002, 43, 2015-20.	3.3	41
141	The cellular fate of mutant rhodopsin: quality control, degradation and aggresome formation. Journal of Cell Science, 2002, 115, 2907-18.	2.0	262
142	The Cochaperone Murine Stress-Inducible Protein 1: Overexpression, Purification, and Characterization. Protein Expression and Purification, 2001, 21, 462-469.	1.3	14
143	Identification and characterization of a human mitochondrial homologue of the bacterial co-chaperone GrpE. Gene, 2001, 267, 125-134.	2.2	24
144	Unfolding retinal dystrophies: a role for molecular chaperones?. Trends in Molecular Medicine, 2001, 7, 414-421.	6.7	99

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145	In vitro studies show that Hsp70 can be released by glia and that exogenous Hsp70 can enhance neuronal stress tolerance. Brain Research, 2001, 914, 66-73.	2.2	306
146	Analysis of the levels of conservation of the J domain among the various types of DnaJ-like proteins. Cell Stress and Chaperones, 2000, 5, 347.	2.9	74
147	Interaction of the human DnaJ homologue, HSJ1b with the 90 kDa heat shock protein, Hsp90. Life Sciences, 2000, 67, 1455-1465.	4.3	16
148	Domain Requirements of DnaJ-like (Hsp40) Molecular Chaperones in the Activation of a Steroid Hormone Receptor. Journal of Biological Chemistry, 1999, 274, 34045-34052.	3.4	43
149	Structure, function and evolution of DnaJ: conservation and adaptation of chaperone function. Cell Stress and Chaperones, 1998, 3, 28.	2.9	528
150	Inhibition of hsc70-catalysed clathrin uncoating by HSJ1 proteins. Biochemical Journal, 1996, 319, 103-108.	3.7	33
151	Regulation of 70â€kDa Heatâ€Shockâ€Protein ATPase Activity and Substrate Binding by Human DnaJâ€Like Proteins, HSJ1a and HSJ1b. FEBS Journal, 1994, 226, 99-107.	0.2	24
152	Regulation of 70-kDa Heat-Shock-Protein ATPase Activity and Substrate Binding by Human DnaJ-Like Proteins, HSJ1a and HSJ1b. FEBS Journal, 1994, 226, 99-107.	0.2	70
153	Expression of alternatively-spliced glutamate receptors in human hippocampus. European Journal of Pharmacology, 1993, 244, 89-92.	2.6	6
154	Apolipoprotein E-∈ A allele and Alzheimer's disease. Lancet, The, 1993, 342, 1308-1309.	13.7	51
155	Isolation of cDNAs coding for epitopes shared by microtubule-associated proteins and neurofibrillary tangles in Alzheimer's disease. FEBS Letters, 1987, 226, 28-32.	2.8	11
156	Intravitreal Sepofarsen for Leber Congenital Amaurosis Type 10 (LCA10). SSRN Electronic Journal, 0, , .	0.4	0