

Table 2

Proteins identified from spots detected only macular retina.

Proteins from macular unique spots by LC-MS/MS and listed.

^(a)Spot ID corresponds to the numbers on gel images in figure 1 - 4.

^(b)Accession no. corresponds to UniProtKB/Swiss-Prot database (Release 48.8).

^(c)MW and pI are theoretical scores by Bioworks ver.3.1.

*Oxidation of methionine.

Spot no. ^(a)	Protein name	Database Accession no. ^(b)	MW (kDa) ^(c)	pI ^(c)	Sequence coverage (%)	Precursor ion MH+	Charge	XC	Residue	Sequence
M1	Pyruvate kinase, isozymes M1/M2	P14618	32.7	4.69	18.11	1198.35	2	3.10	32-42	LDIDSPITAR
						1360.53	2	4.43	43-55	NTGIICTIGPASR
						1194.43	2	2.70	56-65	SVETLKEM*IK
						914.08	2	2.50	106-114	PVAVALDTK
						1119.25	2	2.86	125-135	GSGTAEVELKK
						1463.70	2	4.32	173-185	IYVDDGLISLQVK
						1780.91	2	4.61	188-205	GADFLVTEVENGGSLGSK
M2	Tropomyosin 1 alpha chain	P09493	32.7	4.69	13.73	1142.28	2	2.96	294-304	GDLGIEIPAEK
						1400.56	2	2.95	91-101	RIQLVEEELDR
						1728.89	2	3.24	92-105	IQLVEEELDRAQER
						1315.54	2	3.69	168-178	KLVIIESDLER
						1672.86	2	2.58	169-182	LVIIESDLERAEEER
						1121.21	2	2.35	190-198	CAELEEEELK
						M2	Heterogeneous nuclear ribonucleoproteins C1/C2	P07910	33.7	4.95
1330.47	2	4.35	51-61	GFAFVQYVNER						
1700.00	2	4.50	74-89	M*IAGQVLDINLAAEPK						
1416.60	2	3.49	205-216	QKVDSLLENLEK						
M3	Transaldolase	P37837	37.5	6.36	16.62	1051.20	2	2.82	11-19	M*ESALDQLK
						1792.15	2	4.16	82-97	NAIDKLFVLFGAEILK
						1269.45	2	2.86	111-121	LSFDKAM*VAR
						1234.34	2	2.50	205-215	SYEPLDPGVK
						998.18	2	2.78	231-239	TIVM*GASFR
M3	3'(2'),5'-bisphosphate nucleotidase 1	O95861	33.4	5.46	10.06	1151.34	2	3.98	11-21	LVSAYSIAQK
						1243.43	2	4.16	29-40	VIAEGDLGIVEK
						914.13	2	2.84	225-232	IIQLIEGK
M4	Poly(rC)-binding protein 1	Q15365	37.5	6.66	16.85	1289.37	2	3.24	47-57	INISEGNCPER
						925.13	2	2.25	71-78	AFAM*IIDK

M5	Crk-like protein	P46109	33.8	6.26	7.59	1087.17	2	2.54	315-325	IANPVEGSSGR
M6	Heterogeneous nuclear ribonucleoproteins A2/B1	P22626	37.4	8.97	4.53	1229.45 1318.33	2	3.50	254-265	TALALEVGVYK
M7	Heterogeneous nuclear ribonucleoproteins A2/B1	P22626	37.4	8.97	11.90	1800.00	2	2.79	293-303	IFDPNPDENE
M8	Voltage-dependent anion-selective channel protein 2	P45660	38.1	6.32	9.80	1928.18 1189.34 1696.80	2	4.85	23-38	LFIGGLSFETTESLR
M9	Voltage-dependent anion-selective channel protein 1	P21796	30.6	8.63	7.80	941.03 1722.92 1017.20	2	4.47	22-38	KLPIGGLSFETTESLR
M10	Voltage-dependent anion-selective channel protein 1	P21796	30.6	8.63	15.60	1415.92 1031.23	2	3.72	138-147	IDTIEIITDR
M11	Endoplasmic reticulum protein ERp29	P30040	29.0	6.77	27.99	1860.99 1529.68 1214.35	2	3.17	154-168	GFGFVTFDHDHPVDK
M12	Guanylate kinase	Q16774	21.6	6.11	13.27	1415.92 1031.23	2	2.03	224-235	YOIDPDAFSAK
M13	Guanylate kinase	Q16774	21.6	6.11	19.39	1860.99 1529.68 1214.35	2	3.29	256-265	LTLAALDGGK

1087.17 2 2.54 315-325 IANPVEGSSGR
1229.45 2 3.50 254-265 TALALEVGVYK
1318.33 2 2.79 293-303 IFDPNPDENE
1800.00 2 4.85 23-38 LFIGGLSFETTESLR
1928.18 2 4.47 22-38 KLPIGGLSFETTESLR
1189.34 2 3.72 138-147 IDTIEIITDR
1696.80 2 3.17 154-168 GFGFVTFDHDHPVDK
941.03 2 2.25 193-200 NNFAVGYR
1722.92 2 3.22 263-278 VNNSSLIGVYGTQLR
1017.20 2 2.81 283-292 LTLAALDGGK
1415.92 2 2.03 224-235 YOIDPDAFSAK
1031.23 2 3.29 256-265 LTLAALDGGK
1860.99 2 5.74 34-62 SENGLFTSSGGSANTETTK
1529.68 2 3.53 96-109 LITDSSFSPNTGKK
1214.35 2 3.32 163-173 VTOSNFAVGYK
1325.54 2 2.86 37-48 GALPLDTVIFYK
1248.32 2 2.61 60-69 FDTQYVGEK
1321.51 2 2.43 113-122 ESYVYVYLFYR
1609.72 2 3.64 123-137 DGFENPVPYTGAVK
1725.86 2 4.72 209-223 ILDGEDFPASEMVTR
1135.34 2 3.65 244-253 SLMLTAFQK
1069.24 2 2.84 5-16 PWLSPSGAGK
1675.78 2 3.85 44-57 PGEENGKQYVYVTR
1069.24 2 3.39 5-16 PWLSPSGAGK
1675.78 2 4.23 44-57 PGEENGKQYVYVTR
1387.60 9 66.107

M15	Fatty acid-binding protein, epidermal	Q01469	15.0	6.80	21.64	1674.84	2	4.65	81-96	TVEEAENIAVTSGVVR
						928.11	2	2.22	24-32	ELGVGIALR
						1272.38	2	2.94	61-71	TTQFSCTLGEK
						1026.04	2	2.25	72-80	FEETTADGR
M16	Arrestin-C	P36575	42.8	5.53	14.95	1402.72	2	3.74	48-58	KLFVM*LTCAFR
						1654.85	2	4.47	59-72	YGRDDLEVIGLTFR
						1654.75	2	3.41	145-158	SFCAENPEETVSKR
						1952.20	2	4.11	167-185	KVQFAPPEAGPGPSAQTIR
M17	Arrestin-C	P36575	42.8	5.53	4.64	1824.03	2	4.66	168-185	VQFAPPEAGPGPSAQTIR
M18	Isocitrate dehydrogenase [NAD] subunit alpha	P50213	39.6	6.46	22.40	1607.90	2	2.90	101-115	TPIAAGHPSM*NLLLR
						1392.58	2	3.70	135-146	TPYTDVNIVTIR
						975.13	2	2.74	170-178	LITEGASKR
						1217.36	2	3.94	179-188	IAEFAFEYAR
						1055.23	2	2.38	206-214	M*SDGLFLQK
						1896.29	2	4.58	300-316	DM*ANPTALLLSAVM*M*LR
						1124.30	2	3.03	327-336	IEAACFATIK
M18	Transaldolase	P37837	37.5	6.36	15.73	1051.20	2	2.54	11-19	M*ESALDQLK
						1250.55	2	2.65	87-97	LFVLFGAEILK
						1234.34	2	2.34	205-215	SYEPLDPGVK
						998.18	2	2.35	231-239	TIVM*GASFR
						1393.60	2	3.77	246-258	ALAGCDFLTISPK
M19	Tropomyosin 1 alpha chain	P09493	32.7	4.69	14.79	1885.07	3	4.05	91-105	RIQLVEEELDRAQER
						1977.23	3	5.13	134-149	AQKDEEKM*EIQEIQLK
						1315.54	2	3.66	168-178	KLVIIESDLER
M19	Heterogeneous nuclear ribonucleoproteins C1/C2	P07910	33.7	4.95	12.75	1317.60	2	3.87	18-29	VFIGNLNTLVVK
						1330.47	2	3.69	51-61	GFAFVQYVNER
						1700.00	2	5.06	74-89	M*IAGQVLDINLAAEPK
M20	Pyruvate dehydrogenase E1 component beta subunit	P11177	39.2	6.20	31.75					

						1803.01	2	3.07	53-68	VFLLGEEVAQYDGAYK
						1845.16	2	4.59	130-145	TYYM*SGGLQPPIVFR
						1352.49	2	2.48	259-269	EGVECEVINM*R
						1901.22	2	3.37	270-285	TIRPM*DM*ETIEASVM*K
						1765.03	2	4.39	309-324	IM*EGPAFNFLDAPAVR
						1265.46	2	3.43	325-336	VTGADVPM*PYAK
						1256.43	2	3.25	337-347	ILEDNSIPQVK
M21	Glucose-6-phosphate 1-dehydrogenase	P11413	59.1	6.44	5.84					
						1174.34	2	2.75	182-191	LSNHISLFR
						1265.53	2	2.37	205-214	EM*VQNLM*VLR
						1192.30	2	2.39	498-507	VGFAQYEGTYK
M22	Glucose-6-phosphate 1-dehydrogenase	P11413	59.1	6.44	9.73					
						1139.33	2	2.58	95-103	LKLEDFAR
						1174.34	2	3.07	182-191	LSNHISLFR
						1265.53	2	2.35	205-214	EM*VQNLM*VLR
						1274.41	2	3.75	246-256	GGYFDEFGIIR
						1192.30	2	2.42	498-507	VGFAQYEGTYK
M23	Glucose-6-phosphate 1-dehydrogenase	P11413	59.1	6.44	12.65					
						1011.16	2	2.36	81-88	KQSEPFK
						1174.34	2	2.91	182-191	LSNHISLFR
						1265.53	2	2.24	205-214	EM*VQNLM*VLR
						1274.41	2	3.44	246-256	GGYFDEFGIIR
						1944.13	3	3.81	370-385	LQFHDVAGDIFHQQCK
						1192.30	2	2.72	498-507	VGFAQYEGTYK
M24	26S proteasome non-ATPase regulatory subunit 11	O00231	47.3	6.09	23.04					
						1517.58	2	4.24	33-45	DIQENDEEAVQVK
						1401.63	2	3.41	46-58	EQSILELGSLLAK
						1158.33	2	3.13	59-70	TGQAAELGGLLK
						1324.55	2	3.14	71-81	YVRPFLNSISK
						1086.26	2	2.30	132-140	LVSIFYDTK
						1342.61	2	3.28	163-174	ALLVEVQLLESK
						1731.99	2	4.86	258-273	IM*LNTPEDVQALVSGK
						1267.50	2	3.15	344-354	VQIEHISLIK
M25	Elongation factor Tu	P49411	49.5	7.26	10.62					
						1674.84	2	4.58	105-120	GITINAAHVFYSTAAR

					1186.30	2	3.31	316-327	AEAGDNLGALVR
					1150.35	2	3.50	352-361	VEAQVYILSK
M26	Elongation factor Tu	P49411	49.5	7.26	4.87				
					1186.30	2	3.66	316-327	AEAGDNLGALVR
					1150.35	2	3.14	352-361	VEAQVYILSK
M27	Alpha-actinin	P61163	42.6	6.19	8.78				
					1868.04	2	4.98	239-255	AQYYLPDGSTIEIGPSR
					1684.96	2	3.94	293-308	TLFSNIVLGGSTLTK
M28	Heterogeneous nuclear ribonucleoproteins C1/C2	P07910	33.7	4.95	16.01				
					1317.60	2	3.56	18-29	VFIGNLNTLVVK
					1124.27	2	3.16	30-39	KSDVEAIFSK
					1330.47	2	4.23	51-61	GFAFVQYVNER
					1700.00	2	4.11	74-89	M*AGQVLDINLAAEPK
M29	Heterogeneous nuclear ribonucleoproteins C1/C2	P07910	33.7	4.95	22.55				
					1317.60	2	3.72	18-29	VFIGNLNTLVVK
					996.10	2	2.73	31-39	SDVEAIFSK
					1330.47	2	4.45	51-61	GFAFVQYVNER
					1700.00	2	5.44	74-89	M*AGQVLDINLAAEPK
					1229.45	2	3.27	188-198	LKGDDLQAIKK
					1160.30	2	3.51	207-216	VDLLENLEK
M30	Heterogeneous nuclear ribonucleoprotein H3	P31942	36.9	6.37	11.27				
					1272.39	2	3.88	56-67	STGEAFVQFASK
					1920.12	2	4.95	206-222	ATENDIANFFSPLNPIR
					1053.15	2	2.76	223-232	VHIDIGADGR
M31	Voltage-dependent anion-selective channel protein 1	P21796	30.6	8.63	19.50				
					1960.99	2	4.52	34-52	SENGLEFTSSGSANTETTK
					1401.50	2	4.13	96-108	LTFDSSFSPNTGK
					1214.35	2	3.33	163-173	VTQSNFAVGYK
					1415.52	2	3.13	224-235	YQIDPDACFSAK
M31	Esterase D	P10768	31.5	6.54	4.61				
					1402.53	2	4.63	186-198	KAFSGYLGTDQSK
M32	Pyruvate kinase, isozymes M1/M2	P14618	57.8	7.95	33.21				
					1198.35	2	3.32	32-42	LDIDSPPIAR
					1360.53	2	3.86	43-55	NTGIICTIGPASR
					1885.03	3	4.35	73-88	LNFSHGTHEYHAETIK

					914.08	2	2.85	106-114	PVAVALDTK
					1214.37	2	2.87	141-150	ITLDNAYM*EK
					1463.70	2	4.50	173-185	IYVDDGLISLQVK
					1780.91	2	6.10	188-205	GADFLVTEVENGGSLGSK
					1766.03	2	4.67	206-223	KGVNLPGAAVDLPVSEK
					1682.88	2	4.25	279-293	FDEILEASDGIM*VAR
					1142.28	2	3.10	294-304	GDLGIEIPA EK
					1020.12	2	2.56	367-375	GDYPLEAVR
					996.17	2	2.76	489-497	VNFAM*NVGK
					1084.19	2	3.01	516-525	PGSGFTNTM*R
M33	Pyruvate kinase, isozymes M1/M2	P14618	57.8	7.95	29.43				
					1360.53	2	4.10	43-55	NTGIICTIGPASR
					1571.71	2	4.07	92-105	TATESFASDPILYR
					914.08	2	2.69	106-114	PVAVALDTK
					1214.37	2	3.10	141-150	ITLDNAYM*EK
					1463.70	2	4.43	173-185	IYVDDGLISLQVK
					1780.91	2	5.94	188-205	GADFLVTEVENGGSLGSK
					1637.86	2	4.16	207-223	GVNLPGAAVDLPVSEK
					954.02	2	2.69	270-277	IENHEGVR
					1682.88	2	3.17	279-293	FDEILEASDGIM*VAR
					1142.28	2	3.04	294-304	GDLGIEIPA EK
					1020.12	2	2.57	367-375	GDYPLEAVR
					996.17	2	2.90	489-497	VNFAM*NVGK
					1084.19	2	2.89	516-525	PGSGFTNTM*R
M34	Aspartate aminotransferase	P17174	46.1	6.57	8.74				
					1357.54	2	3.39	86-98	LALGDDSPALKEK
					1428.62	2	4.00	99-113	RVGGVQSLGGTGALR
					1013.09	2	2.45	259-266	NFGLYNER
M35	Heterogeneous nuclear ribonucleoproteins A2/B1	P22626	37.4	8.97	9.92				
					1058.21	2	2.21	4-12	TLETVPLER
					1800.00	2	4.67	23-38	LFIGGLSFETTEESLR
					1189.34	2	2.67	138-147	IDTIEITDR
M36	Heterogeneous nuclear ribonucleoproteins A2/B1	P22626	37.4	8.97	9.92				
					1058.21	2	2.28	4-12	TLETVPLER
					1800.00	2	4.67	23-38	LFIGGLSFETTEESLR

M37	Heterogeneous nuclear ribonucleoproteins A2/B1	P22626	37.4	8.97	7.37	1800.00	2	4.40	23-38	LFIGGLSFETTEESLR
						1189.34	2	2.81	138-147	IDTIEITDR
M38	Heterogeneous nuclear ribonucleoproteins A2/B1	P22626	37.4	8.97	7.37	1800.00	2	4.06	23-38	LFIGGLSFETTEESLR
						1189.34	2	2.99	138-147	IDTIEITDR
M39	Phosphoglycerate mutase 1	P18669	28.7	6.75	8.30	1060.19	2	2.48	90-99	HYGGLTGLNK
						1151.34	2	3.16	180-190	VLIAAHGNSLR
M40	Superoxide dismutase [Mn]	P04179	24.7	8.35	10.36	1425.66	2	3.68	76-89	GDVTAQIALQPALK
						1029.22	2	2.59	115-123	GELLEAIKR

(Neurobiology of Disease)

Expression of Mutated Optineurin Leads to Normal Tension Glaucoma in Mice by the
Disruption of Optineurin-Rab8 Interaction

Abbreviated title: Mutated optineurin leads to normal tension glaucoma in mice

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ABSTRACT

Glaucoma is one of the leading causes of blindness, affecting 70 million people worldwide. Glaucoma is characterized by a progressive loss of retinal ganglion cells and is often associated with elevated intraocular pressure (IOP). However, patients with normal tension glaucoma (NTG), a subtype of primary open angle glaucoma (POAG), are affected without IOP elevation. Molecular pathways leading to the pathology of the disease are still unclear mainly due to the lack of animal models. Here, we describe the first animal model of NTG based on the same gene mutation found in NTG patients. The transgenic mice over expressing E50K mutation of optineurin (*OPTN*) developed phenotype which mimics the clinical features of NTG patients including degeneration of the retinal ganglion cells at normal IOP. We demonstrate that the E50K mutation in *OPTN* disrupts the interaction between *OPTN* and Rab8, a protein known to regulate vesicle transport from Golgi to plasma membrane. Wild-type *OPTN* and active GTP-bound form of Rab8 complex were localized adjacent to the Golgi complex. These data provide new information about the initial steps in the development of NTG at the molecular level and a new animal model of NTG suitable for therapeutic development.

INTRODUCTION

Glaucoma is characterized by progressive loss of retinal ganglion cells (RGCs), degeneration of axons in the optic nerve, and visual field defects. POAG is one of the major causes of irreversible blindness leading to vision loss in about 4.5 million people and accounting for 12% of all global blindness (Quigley et al., 1996; 2006). POAG is often associated with elevated IOP, which is one of main risk factors in glaucoma. However, characteristic degenerative changes in the retina and optic nerve, as well as visual field loss, may occur even in the absence of elevated IOP in a subtype of POAG which is called NTG. Recent epidemiological study in Tajimi city, have resulted with more than 90% of POAG cases diagnosed as NTG (Iwase et al., 2004).

At present, at least 24 different genetic loci have been linked to various forms of glaucoma and over the last decade, four genes, *myocilin*, *cytochrome P4501B1*, *OPTN*, and *WDR36* have been identified as glaucoma-associated genes (Stone et al., 1997; Stoilov et al., 1997; Rezaie et al., 2002; Monemi et al., 2005). *OPTN* was the first identified gene mutation in which can lead to NTG. The *OPTN* gene contains an initial three 3 non-coding exons followed by 13 exons encoding a protein with a length of 577 amino acids. The *OPTN* mutation at nucleotide 458 (G>A) leading to substitution of glutamic acid by lysine at amino acid 50 (E50K) is tightly linked to patients with a severe NTG phenotype. While this mutation has never been reported

in the normal population (Rezaie et al., 2002; Aung et al., 2005; Alward et al., 2003), mutations in the *OPTN* gene have been observed in 13.5% of NTG families.

Chalasanani et al. (2007) demonstrated that the *OPTN* E50K mutation selectively induces RGC death. *OPTN* has been shown to interact with number of proteins including huntingtin (Faber et al., 1998), transcription factor IIIA (Moreland et al., 2000), RAB8 (Hattula and Peranen, 2000; Park et al., 2006), myosin VI (Sahlender et al., 2005), FOS (Miyamoto-Sato et al., 2005), ring finger protein 11 (Colland et al., 2004), and metabotropic glutamate receptor 1-a (Anborgh et al., 2005).

The molecular pathways leading to NTG from a single gene mutation still remain unclear mainly due to the difficulty of working with patient's eyes at the molecular level and lack of animal models with this particular mutation. Recently, it has been reported that the glutamate transporter-deficient mice exhibit an NTG-like phenotype (Harada et al., 2007). However to date, no NTG animal models have been developed based on the gene mutation found in patients with NTG.

In this paper we show that interaction of *OPTN* with the GTP-form of Rab8, which occurs adjacent to the Golgi, is significantly reduced by the *OPTN* E50K mutation. Transgenic mice over expressing this mutant show progressive RGC loss and excavation of the optic nerve head without detectable changes in IOP similar to the phenotype observed in NTG patients. These mice represent the first *OPTN*-based

animal model of NTG.

MATERIALS AND METHODS

Development of transgenic mice over expressing mutant OPTN

Total RNA was extracted from a fresh C57BL/6N mouse brain tissue using TRIzol (Invitrogen, Carlsbad, CA) and reverse-transcribed into first-strand cDNA using oligo-dT adaptor primer and SuperScript First-Strand Synthesis System for RT-PCR (Invitrogen). *OPTN* cDNA was amplified by PCR using oligonucleotides 5'-cgggaattccgatgtcccatcaacctctgag-3' and 5'-cgggaattccgtcaaatgatgcagtcctcatca-3' as primers. The amplified DNA fragment was purified using a MinElute gel extraction kit (Qiagen, Hilden, Germany), ligated into pBluescriptII (KS-) (Agilent Technologies, Santa Clara, CA) and sequenced using the M13 primers and ABI PRISM 3130 (Applied Biosystems, Foster City, CA). Site-directed mutagenesis was carried out to produce cDNA corresponding to the deletion of the E50K mutation, the first leucine zipper (1st LZ del), deletion of the second leucine zipper (2nd LZ del) and the H486R mutation. The following primers were used: 5'-cagctcaaactcaactccgg -3' and 5'-atgctccacttctgctcca -3' for 1st LZ del , 5'-aaatgaaggaactcctggtaagaaccaccagctgaaagaa-3' and 5'-ttcttcagctgggtggttctaaccaggagttccttcattt-3' for E50K, 5'-gagaccatggccgtctc-3' and

5'-caacatctgtccacctttctg-3' for 2nd LZ del, and
5'-gcagcaagagagaagattcgtgaagaaaaggagcagc-3' and
5'-gctgctcctttctcacgaatcttctcttctgctgc-3' for H486R. The identities of all clones were confirmed by sequencing. Plasmids were digested with EcoRI, purified by agarose gel electrophoresis, and recovered using the MinElute gel extraction kit according to the manufacturer's protocol. The cDNA inserts were ligated into EcoRI digested pCMVHA vector (Takara Bio USA, Madison, WI). HA-tagged *OPTNs* were amplified by PCR using oligonucleotides 5'- ccgctcgagcgcaccatgatgtaccatacagatgtcc-3' and 5'- ccgctcgagcggcctcaaatgatgcagtcctca-3' as primers. The amplified DNA fragments were purified using a MinElute gel extraction kit (Qiagen), ligated into the pCAGGS vector and sequenced as above. cDNA inserts were released from the pCAGGS vector using Sall and BamHI. These restriction fragments were injected into pronuclear stage BDF1/C57BL6N embryos and transgenic mice were generated at PhoenixBio Co., Ltd (Tochigi, Japan). Offspring were screened for the transgene by isolating genomic DNA from tail biopsies followed by PCR. Primers used for PCR were 5'-ctctagagcctctgctaaccatgt-3' and 5'- ccatggccataagagcgtaa -3'. All experiments with mice were performed in accordance with the Association for Research in Vision and Ophthalmology statement for the Use of Animals in Vision Research.

Fundus photography

Mice were anesthetized by aspiration of isoflurane (Mylan, Canonsburg PA). Optic disk imaging was performed as previously described (Coban et al., 2003). Fundus images were obtained using 2 mm gonio lens (Ocular Instrument, Bellevue, WA) and slit lamp (SR-D7, Topcon, Tokyo, Japan) with digital camera (D100, Nikon, Tokyo, Japan).

Light microscopic histopathology of the optic nerve

After deep anesthesia, mouse eyes were dissected and immersed in Davidson solution fixative overnight at 4°C. The eyes were embedded in paraffin and sectioned at 5 µm thickness along the vertical meridian through the optic nerve head. After deparaffinization, and rehydration, sections were stained with hematoxylin and eosin (HE staining).

Electroretinogram (ERG)

Mice (15 month) were anesthetized by intraperitoneal injection of a mixture of xylazine (10 mg/kg) and ketamine (25 mg/kg). Pupils were dilated with 0.5% phenylephrine hydrochloride and 0.5% tropicamide. ERGs were recorded by Mayo Co. (Nagoya, Japan). Standard Flash ERG was obtained using flash intensity 2.50cd·

s/m² with a single flash. To isolate cone response, eyes were light adapted for 10 minutes with a 25cd/m² adapting field. Cone ERGs were obtained using flash intensity 2.50cd·s/m² presented at 0.5Hz with 50 flashes averaged for this intensity. Electrical signals were amplified with 8,300 gain and a bandwidth of 0.5 to 500Hz, electrical signals were digitized at a 10kHz rate using Evoked Potential/EMG Measuring System (Neuropack μ MEB-9102, Nihon Kohden Corporation, Tokyo, Japan).

Immunohistochemistry

The eyes were sectioned at 5 μ m thickness along the vertical meridian through the optic nerve head. After deparaffinization and rehydration, the tissue sections were treated with Target Retrieval Solution (DakoCytomation, Denmark). The sections were incubated with blocking solution for 1 h followed by overnight incubation with primary antibody against HA tag (1:500 dilution; Sigma-Aldrich, St. Louis, MO), tublin β III isoform (1:100 dilution; Millipore, Billerica, MA), OPTN (1:500 dilution; kind gift from Mansoor Sarfarazi), or Calretinin (1:500 dilution; Sigma) in PBS containing 1% BSA at 4°C. Slides were washed in PBS and then incubated with Alexa 488 or Alexa 568 (1:500 dilution; Invitrogen) conjugated secondary antibody and with 4', 6'-diamidino-2-phenylindole (DAPI) for nuclear staining for 1 h at room temperature.

The stained tissues were examined using confocal fluorescence laser microscope (Radiance 2000, Bio-Rad Laboratories, Hercules, CA). For negative control of the immunohistochemical staining, the sections were incubated with blocking solution without primary antibody (data not shown).

Whole-mount immunostaining

The whole-mount immunostaining was performed essentially as described (Jakobs et al., 2005). Anterior parts were dissected from enucleated eyes. The posterior parts were fixed in 4% PFA/PBS for 2 hours on ice and then incubated with the anti-SMI32 (1:200 dilution; Sternberger Monoclonals, Baltimore, MD), anti-GFAP (1:200 dilution; Millipore) or anti-active caspase-3 (1:200 dilution; Imgenex, San Diego, CA) antibody for 7 days at 4°C. Slides were washed in PBS containing 0.1% Triton X-100 and then incubated with Alexa 488 or Alexa 568 (1:500 dilution; Invitrogen) conjugated secondary antibody and with 4', 6' -diamidino-2-phenylindole (DAPI) for nuclear staining for 2 days at 4°C. The retinas were then mounted with Vectashield (Vector Laboratories, Burlingame, CA) and evaluated on a confocal microscope.

Measurement of intraocular pressure (IOP)

The average IOP for each genotype was recorded. IOP was measured using an

impact-rebound tonometer (Colonial Medical Supply, Franconia, NH) and optical interferometry tonometer (FISO Technologies, Quebec, Canada) for mice of each genotype as described (Senatorov et al., 2006). Using the rebound tonometer, we were able to measure IOP in awake and non-sedated mice of various ages, while optical interferometry tonometry was performed on anesthetized animals. Measurement of IOP was always performed in the morning between 10 and 12 a. m. The numbers of mice successfully assessed for each genotype and age were 18 weeks and 12 month.

Measurement of OPTN-Rab8 interaction

OPTN-Rab8 interaction analysis was performed using *CoralHue*[®] Fluo-chase Kit (MBL, Tokyo, Japan). Based on the instruction manual, we constructed *OPTN* wild type, *OPTN* E50K, and Rab8 wild type, Rab8 GDP-bound (T22N) inactive form, and Rab8 GTP-bound (Q67L) active form recombinant fusion protein plasmids with both N-terminal and C-terminal fluorescence protein (Fig. 7A). RGC-5 and COS1 cells were transfected by each pairs of the plasmid mixtures using Fugene HD (Roche). 48 h after transfection, the medium was replaced to phosphate buffered saline (PBS) and the cells were observed with inverted microscope (Eclipse TE300, Nikon). To observe OPTN-Rab8 complex, cells were fixed 48 h after transfection with 4%