Deciphering the Structure and Function of Als2cr4 in the Mouse Retina

Freddi I. Zuniga1,2 and Cheryl M. Craft1,2,3

PURPOSE. The role of Als2cr4 (amyotrophic lateral sclerosis 2 [juvenile] chromosome region, candidate 4; also known as hypothetical protein FLJ33282) in the mouse retina was determined by characterizing the molecular structure, cellular interacting partners, and potential biochemical functions. Previous in situ hybridization and gene expression profiles show that the mRNAs encoding Als2cr4 are abundant in the eye, hippocampus, cerebellum, and olfactory bulb.

METHODS. From predicted antigenic epitopes of Als2cr4, two novel antibodies were developed to examine protein expression and morphologic localization in retinas from light-adapted and dark-adapted mice by immunohistochemistry, immunoblot analysis, and immunoelectron microscopy, and then immunoprecipitation was performed to identify interacting proteins by mass spectrometry.

RESULTS. Peptide antibodies with Als2cr4 antigenic epitopes from either the amino- or carboxyl terminus were characterized with Als2cr4 recombinant proteins and peptide competition assays. Als2cr4 is a 45kDa insoluble protein, highly enriched in retina, and localizes to photoreceptor outer segments, ciliary complex, and horizontal cells in the outer plexiform layer. Immunoelectron microscopy for Als2cr4 verified its expression in the discs of photoreceptor outer segments. Immunoprecipitation and mass spectrometry identified eight potential interacting partners: vimentin, actin, myosin Va, myosin VI, myosin X, myosin XIV, kinesin 1, Als2cr4, and lamin B-1.

CONCLUSIONS. Als2cr4 is a novel protein, with a probable tetrarospan-like membrane structure, that is localized in photoreceptors and in the postsynaptic outer plexiform layer and that interacts with cytoskeletal proteins. Als2cr4 may be involved in membrane transport between the photoreceptor inner and outer segments and may be a key component in maintaining the structural integrity of the outer segment. (Invest Ophtalmol Vis Sci. 2010;51:4407–4415) DOI:10.1167/iovs.10-5251

The vertebrate retina is an intricate organ consisting of five distinct types of neurons: photoreceptors, horizontal cells, bipolar cells, amacrine cells, and ganglion cells. These specialized cell types process information in the form of light quanta and relay information to the lateral geniculate nucleus in the brain and ultimately to the primary visual cortex. Photoreceptors are the polarized, light-sensitive cell types responsible for initiating the phototransduction cascade in the mammalian retina and are of two types: rods and cones, with the predominant ones being rods. Although photoreceptors share similar cellular mechanisms for the activation and deactivation of phototransduction, they respond to various intensities of light and show distinct in the wavelengths of light absorbed by the visual pigments.

Although much is known about the visual process, a better understanding of phototransduction is essential, since many visual impairments develop as a direct result of mutations in the genes encoding the proteins essential for phototransduction. One such protein is arrestin 1 (ARR1), which, when defective, leads to a form of retinitis pigmentosa known as Oguchi disease, which has congenital stationary night blindness phenotype. Arrestins downregulate activated, phosphorylated G-protein-coupled receptors and include four subtypes: ARR1 and arrestin-4 (ARR4, cone or X-arrestin) and the ubiquitously expressed β-adrenergic arrestin-1 and -2 (ARR2 and -3). To identify and characterize potential functional partners for ARR4, we screened a mouse retinal cDNA yeast two-hybrid (Y2H) library by using ARR4 as bait (Zuniga FI, et al. IOVS 2007;48:ARVO E-Abstract 601; Zuniga FI, et al. IOVS 2009;50: ARVO E-Abstract 5453). From the list of candidate proteins, Als2cr4 (NM_001037812, hypothetical protein FLJ33282; accession numbers are all GenBank; http://www.ncbi.nlm.nih.gov/Genbank; National Center for Biotechnology Information [NCBI], Bethesda, MD) was further investigated. Als2cr4 was first identified in Mus musculus in 2005 by the Laboratory for Genome Exploration Research Group at the RIKEN Genomic Sciences Centre (Kanagawa, Japan). The official gene symbol is Als2, corresponding to mouse amyotrophic lateral sclerosis (ALS) 2 (juvenile) chromosome region, candidate 4. As indicated by the name, this gene was in a candidate region for the neurodegenerative disease ALS; however, it was found that defects in ALS2 and its encoded protein, alsin, are responsible for neurotoxicity by the familial ALS-linked mutant Cu/Zn-superoxide dismutase (FOSD1). In the mouse, Als2cr4 is located on chromosome 1 and consists of two isoforms (1 [NP_001028621] and 2 [NP_001032901]), sharing more than 95% identity. The divergent sequence is located at the extreme amino terminus: amino acids (AA) 1-21. Moreover, Als2cr4 shares approximately 83% identity with its human ortholog located on 2q33.2. The in situ hybridization (St. Jude’s Brain Gene Expression Map www.stjudebgem.org/ provided in the public domain by St. Jude’s Research Hospital, Memphis, TN) gene expression patterns show that the mRNA is highly abundant in the mouse eye from embryonic day (E)15 through...
adulthood. Prominent expression was also noted in the hip- 
campus, cerebellum, and olfactory bulb from postnatal day 
(P)7 through adulthood.

Recent technological advances are now available to aid in 
the study of the proteome of photoreceptors and have led to 
previously unidentified candidates that cause visual dysfunc-

tion. In two retinal proteomic analyses, Als2cr4 was identified 
in mouse sensory ciliary complex (CG1100) and in a rod outer 
segment (OS) preparation.11

We explored the structural and biochemical role that this 
novel protein plays in the retina under different environmental 
lighting conditions and in vitro cultures by using newly de-
veloped antibodies for Als2cr4 with immunoblot (IB) analysis, 
immunohistochemistry (IHC), and immunoelectron micros-
copy (IM-EM). In addition, to differentiate between the role 
of Als2cr4 in rod and cone photoreceptors, we compared retinas 
from C57Bl/6J (wild-type; WT) mice with those of neural retina 
leucine zipper knockout mice (Nrl−/−). Nrl is a key transcrip-
tion factor responsible for the developmental program switch 
of rods to enhanced S-cone expression.12 Using these reagents 
for co-immunoprecipitation (IP), we identified interacting pro-
teins through mass spectrometry.

METHODS

Animals

All procedures involving mice were approved by the Institute for 
Laboratory Animal Research and conducted according to the ARVO 
Statement for the Use of Animals in Ophthalmic and Vision Research. 
Original breeders for the Nrl−/− and Arr4+/− mice were generated and 
generously provided by Anand Swaroop (University of Michigan, 
NIH-NII)12 and Jeannie Chen (University of Southern California; 
USC),13 respectively. The Arr4+/− and Arr4+/−/Arr4+/− double-knock-
out mice were generated in our laboratory.14 The WT (C57Bl/6J and 
129SvJ mixed background) mice were from our breeding colony. All 
animals were born and maintained in the USC vivarium in the Zilkha 
Neurogenetic Institute. They were kept under controlled ambient 
illumination on a 12-hour:12-hour light–dark cycle, with the exception 
of their susceptibility to light-induced degeneration.15

Computer-Generated Primary and 
Two-Dimensional (2-D) Secondary Structure

The Als2cr4 (NP_001032901) predicted translated AA sequence was 
used to identify protein transmembrane helices with version 2 of the 
TMHMM Server (Transmembrane Helices Markov Model; http:// 
www.cbs.dtu.dk/services/TMHMM-2.0; provided in the public domain 
by the Center for Biological Sequence Analysis, the Technological 
University of Denmark, Lyngby, Denmark). These data were input into 
TMRPres2D software (http://bioinformatics.biol.uoa.gr/TMRPres2D/ 
download.jsp) to create a 2-D model of α-helical or β-barrel transmem-
brane proteins.

Antibody Production

The mouse Als2cr4 AA sequence was assessed for hydrophilicity, 
antigenicity, surface accessibility, and predicted secondary structure 
by using Invitrogen’s open source PeptideSelect Online Designer soft-
ware (http://peptideselect.invitrogen.com/peptide/; Invitrogen, Carls-
bad, CA). Candidate sequences were subjected to BLAST queries to 
circumvent cross-reactivity with other proteins (www.ncbi.nlm. 
nih.gov/blast/ provided in the public domain by NCBI, Bethesda, MD).

The Zymed Laboratory (Invitrogen) developed two affinity-purified 
polyclonal antibodies (pAbs) with predicted antigenic epitopes (FLJ-
Frederick and Doric Miller [FLJ-FM] AA 76-88 EAPAPRqQKIRRp; and 
[FLJ-Thomas and Laurene Gray [FLJ-LG] AA 390-403 DVDEHPETG-
KASP) to the amino- and carboxyterminal domains, respectively.

Immunoblot Analysis

The immunoblot analysis using sodium dodecyl sulfate-polyacrylamide 
gel electrophoresis (SDS-PAGE) was modified from previously pub-
lished protocols.16 Briefly, mice were kept in darkness for 24 hours, 
exposed to room light for 10 minutes, and killed or dark adapted (DA) 
on the retina dissected under infrared (IR) light. The retinas were homogenized (Polytron PT1200; Kine-
matica AG, Lucerne, Switzerland) in NP-40 lysis buffer (50 mM Tris-
Cl [pH 7.4], 250 mM NaCl, 5 mM EDTA, 1% NP-40, and 0.02% Na3P, 
plus protease inhibitor cocktail; Halt Protease/Phosphatase Inhibitor Cock-
tail; Thermo Scientific, Rockford, IL) and centrifuged at 13,000 rpm for 
15 minutes at 4°C. The pellet fractions were further disrupted (Micro-
son sonicator, Misonex, NY) in minimum CHAPS lysis buffer (50 mM 
Tris-Cl [pH 7.6], 150 mM NaCl, 10 mM CHAPS, and 0.02% Na3P, plus 
protease/phosphatase inhibitor cocktail) and centrifuged as just de-
scribed. The cell lysates were pooled, and 20 μg of retinal homogenate 
was mixed with Laemmli buffer, boiled for 10 minutes, and resolved on 
10% to 12% SDS-PAGE. Immobilized proteins were stained with Coom-
assie blue or transferred onto PVDF membrane (Immobilon-P PVDF; 
Millipore, Billerica, MA) and reversibly stained (MemCode Reversible 
stain; Thermo Scientific). The membranes were blocked in 5% milk 
for 30 minutes at room temperature (RT), incubated with primary anti-
body overnight, labeled with horseradish peroxidase (HRP)-conju-
gated secondary antibody (Bio-Rad, Hercules, CA), and visualized 
by enhanced chemiluminescence (ECL) detection. Antibodies were used 
at the following concentrations: FLJ-FM pAb, 1:1,000; FLJ-LG pAb, 
1:1,000; anti-GAPDH mAb (G8795; Sigma-Aldrich, St. Louis, MO), 
1:5,000; anti-Arr1 mAb (D9F2), 1:50,000; anti-Arr4 rabbit pAb (mCAR-
LUMI), 1:1,000; anti-myosin Va pAb (sc-17707; Santa Cruz Biotechnol-
ology, Santa Cruz, CA), 1:375; and anti-myosin VI pAb (sc-23,68; Santa 
Cruz), 1:1,500. The secondary antibodies goat anti-rabbit HRP-conju-
gated IgG (170-6515; Bio-Rad) and goat anti-mouse HRP-conjugated IgG 
(172-1011; Bio-Rad) were used at 1:10,000.

Peptide Competition Assay

Samples were prepared as just stated, resolved on 12% SDS-PAGE, 
transferred onto PVDF membrane, reversibly stained (MemCode Re-
versible Protein stain; Thermo Scientific), and cut into equal halves. 
Als2cr4 antibody (FLJ-FM pAb) was preadsorbed with excess FLJ-FM 
peptide in 5% milk overnight at 4°C with gentle agitation. The mem-

branes were probed with FLJ-FM pAb or preadsorbed FLJ-FM pAb (+ 
peptide) overnight, followed by secondary goat anti-rabbit HRP-conju-
gated antibody for ECL detection.

Immunoprecipitation

Retinal lysates from WT or Nrl−/− mice were prepared in either room 
light or in total darkness, as stated earlier. Each procedure was con-
donced in its respective lighting condition throughout the experiment. 
Magnetic beads (Dynal; Invitrogen) were prepared and incubated with 
retinal lysate for 2 hours at 4°C with end-over-end shaking. The pre-
clared supernatant was collected and incubated with 10 μg of pAb 
FLJ-FM or a nonspecific antibody at 4°C overnight. Magnetic protein-G 
beads were added and incubated for 2 hours at 4°C with shaking. The 
beads were gently washed three times with 1× PBS before 5× SDS 
sample loading buffer (2× final concentration) was added and the 
mixture heated at 95°C for 10 minutes. The precipitated proteins were 
resolved on 10% SDS-PAGE and either stained with Coomassie blue for 
mass spectra analysis or transferred to PVDF membranes and subjected 
to IB analysis.

Mass Spectrometry

Selected fragments from the IP identified by pAb FLJ-FM were excised 
from the Coomassie blue-stained gel; in-gel trypsin digestion was 
performed and tryptic fragments were analyzed by liquid chromatog-
raphy with tandem mass spectrometry detection (LC-MS/MS).17 A por-
tion of the digested peptide mixture was also analyzed with a custom-

tide prophet algorithm.20 Protein identifications were accepted if they established at greater than 50.0% probability, as specified by the peptide-protein algorithm.20 Protein identifications were accepted if they could be established at greater than 95.0% probability and contained at least five identified peptides. Protein probabilities were assigned by the protein algorithm.21

**Immunohistochemistry**

One-month-old mice were DA for 24 hours, exposed to room light for 1 hour and euthanatized or kept in total darkness where they were killed and the eyecups dissected under infrared light.22 The eyecups were fixed in 4% paraformaldehyde (PFA) for 2 hours at 4°C, incubated in 30% sucrose overnight, embedded in optimal cutting temperature (OCT; Sakura Finetek, Torrance, CA) embedding medium, and cryosectioned (model CM 1800; Leica-Jung, Wetzlar, Germany) at 7 μm through the optic nerve head; the slides were stored at −80°C. Tissue slides were thawed for 30 minutes at RT, washed twice in 1× PBS-Tween for 5 minutes, and blocked for 30 minutes with blocking buffer (10% normal goat serum, 0.2% Triton X-100, in 1× PBS) in a humidified chamber. The sections were incubated with primary antibody overnight at 4°C in antibody dilution buffer (2% goat serum, 0.2% Triton X-100, in 1× PBS) with gentle agitation. The day after, the slides were washed three times in 1× PBS-Tween for 15 minutes followed by direct protein labeling with Alexa Fluor-conjugated secondary antibodies (Invitrogen), and the nuclei were counterstained (TO-PRO-3, 1:2,500; secondary antibodies: Alexa Fluor-conjugated IgG, 1:500; and anti-Arr1 mAb (D9F2), 1:20,000; anti-Arr4 pAb (Chicken LUMIj), 1:1,000; anti-calbindin-D-28K mAb (C9848; Sigma-Aldrich), 1:2,000; and a nuclear stain TO-PRO-3, 1:2,500.

**Immunoelectron Microscopy**

One-month-old WT mice were light-adapted (LA) or DA and euthanatized, and the eyes were enucleated. The eyecups were fixed in 4% PFA, washed three times (10 minutes) in fixative buffer, and dehydrated in graded alcohol (30%-100%). Eyecups were then infiltrated with various ratios of resin (LR White Medium Grade, EMS, Hatfield, PA) to ethanol, embedded in fresh resin in gelatin capsules, and polymerized under UV light for 48 hours at 4°C. Sections were cut on a microtome (Ultracut E Ultramicrotome; Reichert, Vienna, Austria) with a diamond knife and placed on uncoated 75/500-mesh nickel grids. Grids were incubated in 0.02 M glycine in 1× PBS for 15 minutes, washed four times in PBS, blocked (5% NGS, 1% BSA in PBS) at RT for 15 minutes, and incubated overnight at 4°C in primary antibody. Sections were washed five times in PBS and incubated in goat anti-rabbit 10-nm colloidal gold (1:50) for 2 hours at RT. The grids were washed three times in PBS, washed twice for 3 minutes each in distilled water (DW), fixed for 5 minutes with 1% glutaraldehyde in DW, washed 30 minutes in 5 drops of DW, and stained with aqueous 2% uranyl acetate in darkness. This procedure was followed by a wash in DW, staining for 15 minutes with Reynolds’ lead citrate at RT and a wash in DW. The grids were allowed to dry and fumed with aqueous 2% osmium tetroxide for 30 minutes at RT. The sections were viewed with an electron microscope (100CX, JEOL, Tokyo, Japan), as previously described.23,24

**cDNA Constructs for Als2cr4**

The cDNA encoding mouse Als2cr4 was obtained from total RNA of retinae from Nrl−/− mice by using a first-strand cDNA synthesis kit (SuperScript III; Invitrogen). The complete predicted coding region of Als2cr4 (NM_001037812) was amplified by polymerase chain reaction (PCR) technology with sense, 5'-ATGGGGAAGAGCAGTGTT3', and antisense, 5'-CTATGGAGAGGCTTTGGTT-3', primer pairs. The amplified product was purified and then subcloned into the plasmid vectors pTrcHis-TOPO and pcDNA4/HisMax-TOPO (Invitrogen). The integrity and nucleotide sequence of the constructs was confirmed by automated DNA nucleotide sequencing.

**Cell Culture**

HEK293 (ATCC, Manassas, VA) cells were grown in minimum essential medium (MEM; ATCC), supplemented with 10% bovine serum albumin (BSA) and penicillin/streptomycin antibiotics in 5% CO2 at 37°C until confluent. The cells were plated on four-well culture slides (BD Falcon, Lincoln Park, NJ) at an initial density of 50,000 cells/chamber and...
grown for 24 hours. Plasmid cDNAs were transfected into HEK293 cells (FugeneHD Transfection reagent; Roche, Mannheim Germany) as suggested in the manufacturer’s protocol. The HEK293 cells were transfected with either pcDNA4-Als2cr4-HisMax or empty vector and allowed to incubate for 24 hours before they were washed with ice-cold PBS and fixed in 4% PFA for 15 minutes at 4°C. The cells were blocked with 10% goat serum and 0.2% Triton X-100 in 1/100 PBS for 30 minutes at RT, followed by incubation for 1 hour at RT with primary antibody (diluted in 2% goat serum and 0.2% Triton X-100 in 1/100 PBS). The cells were washed three times in ice-cold PBS and then incubated with the appropriate secondary antibody before being mounted with coverslips and viewed with a laser scanning confocal microscope (Carl Zeiss Meditec) at 63× magnification. The antibodies were used at the following concentrations: FLJ-FM pAb, 1:1,000; anti-6xHis mAb (H1029; Sigma-Aldrich, MO), 1:3,000; and secondary antibody Alexa Fluor-conjugated IgG, 1:500.

RESULTS

Primary and Secondary 2-D Protein Structure of Als2cr4

To elucidate the predicted structure of Als2cr4, the translated AA sequence was subjected to a functional domain query. The NCBI conserved-domain database identified a multidomain conservation between the Als2cr4 isoform 1 and DNA polymerase III subunits gamma and tau.25 The protein coding sequence of Als2cr4 isoform 2 is composed of 1212 nucleotides (nt 74-1285), and the translational product consists of 403 AAs (NP_001032901), resulting in a predicted translated protein of molecular mass of ~45 kDa. Furthermore, the translated coding sequence of mouse Als2cr4 (isoform 2) was aligned with various orthologs (Supplementary Fig. S1, http://www.iovs.org/cgi/content/full/51/9/4407/DC1) and resulted in the following percent identities: 98% (Mus musculus isoform 1), 81% (Homo sapiens isoform a), 82% (Homo sapiens isoform b), 82% (Macaca mulatta), 79% (Bos taurus), and 49% (Danio rerio).

Computer modeling studies (Fig. 1) predict that the Als2cr4-translated protein contains four transmembrane helices (AAs 228-250, 265-287, 300-322, and 351-373), along with a lengthy 226-AA amino terminus and a short 30-AA carboxyl terminal tail, both predicted to be in the cytoplasmic space. In addition to these structural motifs, Als2cr4 contains a short (15 AA) loop linking transmembrane helices 1 and 2, as well as a longer (29-AA) loop linking helices 3 and 4, both in the extracellular space. An additional 13-AA loop is present in the cytoplasmic space linking helices 2 and 3. Moreover, Als2cr4 is predicted to have a tetratricopeptide motif present in the long (160-192-AA) tail. Tetratricopeptides are structural motifs that form scaffolds to mediate protein–protein interaction.26 The prediction that Als2cr4 is a structural membrane protein agrees with our biochemical and morphologic analyses, as we observed Als2cr4 to be present exclusively in the insoluble retina fraction (Supplementary Fig. S2) and to be embedded within the membrane discs of mouse photoreceptor OS.
To confirm the expression of this newly identified protein, we generated polyclonal antibodies to two unique antigenic epitope domains: FLJ-FM AA 76-88 and FLJ-LG AA 390-403. Antibody specificity was assessed by peptide competition assays (Supplementary Fig. S2) and detection of Als2cr4 recombinant proteins (data not shown). In Supplementary Figure S2, FLJ-FM antibody specificity was abolished by using excess FLJ-FM immunizing peptide. Next, Als2cr4 was tested for differential expression in visual knockout mice (Arr1/H11002/H11002, Arr4/H11002/H11002, Arr1/H11002/H11002 Arr4/H11002/H11002 [Arr double knockout], and Nrl/H11002/H11002) compared with control WT and also differences in expression in LA and DA conditions. In control WT retina, IB analysis (Fig. 2) of retinal lysates showed that Als2cr4 is expressed ubiquitously as a 45-kDa protein. Additional experiments with protein homogenates from the Nrl/H11002/H11002 retina produced a doublet of 45 and 52 kDa, respectively, suggesting that Als2cr4 isoform 1 is a cone-enriched isoform below detectable levels in WT retina that contains 3% to 5% cones.27 In addition, the mice from DA environment compared with LA have similar Als2cr4 protein expression levels (Fig. 2). Furthermore, no differences in Als2cr4 expression patterns were observed on either IB analysis (Fig. 2) or immunofluorescent localization (data not shown) when either Arr1 or Arr4 was knocked out. Its absence may be explained by brief interaction between these proteins or by the use of nonoptimal conditions.

**Cellular and Subcellular Immunohistochemical Localization of Als2cr4**

Our immunofluorescence analysis revealed that Als2cr4 is expressed in both rods (Figs. 3, 4, 5) and Nrl/H11002/H11002 cones produced a doublet of 45 and 52 kDa, respectively, suggesting that Als2cr4 isoform 1 is a cone-enriched isoform below detectable levels in WT retina that contains 3% to 5% cones. In addition, the mice from DA environment compared with LA have similar Als2cr4 protein expression levels (Fig. 2). Furthermore, no differences in Als2cr4 expression patterns were observed on either IB analysis (Fig. 2) or immunofluorescent localization (data not shown) when either Arr1 or Arr4 was knocked out. Its absence may be explained by brief interaction between these proteins or by the use of nonoptimal conditions.

**FIGURE 4.** Dual IHC of Als2cr4 and Arr1 in retinas from light- or dark-adapted mice. P30 WT mice were either (A) light- or (B) dark-adapted and killed. The retina sections were dual stained with mAb D9F2 for Arr1 and pAb FLJ-FM for Als2cr4. Secondary antibodies were Alexa Fluor-568 goat anti-mouse IgG (Arr1) and Alexa Fluor-488 goat anti-rabbit IgG (Als2cr4). Nuclei were stained with TO-PRO3 iodide. (A, top) A magnified region of the merged photoreceptor layer in the bottom panel demonstrates that Als2cr4 was present in the connecting cilia. After 1 hour of light exposure, Arr1 localization was exclusive to the OS, whereas Als2cr4 was present in both the OS and connecting cilia. (B) Als2cr4 was localized in the postsynaptic OPL, with no co-localization of Arr1 with Als2cr4 in rod spherules or cone pedicles. Magnification, ×63.

**FIGURE 5.** Dual IHC localization of Als2cr4 and calbindin in horizontal cells. P30 WT mice were either (A) light- or (B) dark-adapted and killed. The retinal sections were dual stained with mAb calbindin and pAb FLJ-FM. Secondary antibodies were Alexa Fluor-568 goat anti-mouse IgG (Calbindin) and Alexa Fluor-488 goat anti-rabbit IgG (Als2cr4). The nuclei were stained with TO-PRO3 iodide. Top: magnified region of the photoreceptor layer. Middle: view of the retina at ×63 magnification. Bottom: magnified region of the OPL. White arrows denote dual immunohistochemical localization of Als2cr4 with calbindin at the OPL. Magnification, ×63.
green) and 6xHis mAb (red). In Figures 7b, 7d, and 7f, the
tern was visualized by dual-IHC labeling with pAb FLJ-FM
recombinant protein is expressed. The Als2cr4 expression pat-
tagged protein, as well as to determine where the subcellular
and physiological effects that were due to introduction of the
transfected into HEK293 cells to study potential morphologic
retinas (Fig. 4), more studies are warranted.

Expression of the His tag was limited to the
discs of photoreceptor OS. Magnification, ×20,000.

(Supplementary Fig. S3). This observation is in agreement
with our IB analysis showing Als2cr4 expression in WT and
Nrf2−/− mice (Fig. 2). As shown in Figures 3, 4, and 5 with
immunohistochemical localization, Als2cr4 localized pre-
dominantly to photoreceptor OS, CC, and the outer plexi-
form layer (OPL). Dual immunohistochemical labeling exper-
iments with a well-characterized monoclonal D9F2 antibody to
Arr1 and pAb-FLJ-FM to Als2cr4 (Fig. 4A) showed that in light,
Arr1 is localized to the rod and cone OS, whereas Als2cr4 is
present both in the photoreceptor OS and in the CC. A higher
magnification of the photoreceptor layer in Figure 4A (top)
clearly shows the limits of the OS, and dotted white lines
indicate the CC where Als2cr4 staining was also observed. The
Als2cr4 immunostaining pattern appeared punctate in the pho-
toreceptor layer and uniform in the OPL. We observed a similar
immunologic staining pattern in the DA (Figs. 5B, 4B) retinas.
Figure 4B shows that in DA WT retina, Arr1 and Als2cr4 do not
co-localize in either rod spherules or cone pedicles, indicating
that Als2cr4 expression is postsynaptic. Furthermore, dual im-
munohistochemical stain labeling experiments (Fig. 5) demon-
strated that Als2cr4 co-localizes with calbindin, a well-charac-
terized marker used to identify horizontal cells. At present,
however, we are not certain whether expression of Als2cr4 is
limited to horizontal cell axons, dendrites, or both.

In other experiments, the photoreceptor ultrastructure
was examined by IM-EM to visualize the subcellular location
of Als2cr4. Figure 6 shows data from LA (Fig. 6A) and DA
(Fig. 6B) WT retinas. IM-EM with pAb FLJ-FM shows colloidal
gold particles present in the discs of photoreceptor OS.
Figure 6 shows rod OS; however, we observed that cone OS
were also labeled in other micrographs (data not shown). In
addition, IM-EM results suggested that more Als2cr4 is ex-
pressed in LA (Fig. 6A) than in DA (Fig. 6B) retinas; how-
ever, IB (Fig. 2) and IHC experiments (Figs. 3, 4, 5) revealed
comparable protein expression levels in both lighting con-
ditions. These experiments should be replicated and the
results analyzed statistically. With regard to the presence of
Als2cr4 in connecting cilia, which has been reported in the
CC proteome and which we showed by IHC in mouse
retinas (Fig. 4), more studies are warranted.

Subsequently, immunohistochemical localization of exoge-
nous Als2cr4 expression and localization in cultured cells was
examined. The 6xHis-tagged cDNA for Als2cr4 was transiently
transfected into HEK293 cells to study potential morphologic
and physiological effects that were due to introduction of the
tagged protein, as well as to determine where the subcellular
recombinant protein is expressed. The Als2cr4 expression pat-
tern was visualized by dual-IHC labeling with pAb FLJ-FM
(green) and 6xHis mAb (red). In Figures 7b, 7d, and 7f, the
transfected cells show poly-histidine-tagged Als2cr4 recruit-
ment to the cytoplasm and plasma membrane. This observation
correlates with our findings that Als2cr4 was present only in
the pellet fractions of retinal homogenates and not in the
soluble fractions (Supplementary Fig. S2). Furthermore, trans-
faction of 6xHis-Als2cr4 did not disrupt or change cellular
morphology. Last, to resolve whether Als2cr4 is recruited to
the plasma membrane because of its primary AA sequence or
whether the localization is an artifact of the epitope tag, a
control pCDNA4/HisMax without a cDNA insert was trans-
fected into the cells. The data confirm (Figs. 7a, 7c, 7e) that
expression of pCDNA4/HisMax without the recombinant
Als2cr4 insert was limited exclusively to the nucleus, thereby
substantiating that the 6xHis-tag alone would not drive expres-
sion of the His tag control to the cytoplasm or the plasma
membrane.

**IP Showing Potential Interacting Partners of Als2cr4**

IP experiments were performed to identify potential bind-
ing partners for Als2cr4 in response to LA and DA conditions.
Supplementary Figure S4 shows DA IP studies that were ana-
lyzed from stained gels with either a control IgG or Als2cr4-
specific pAb-FLJ-FM. Lane 1 represents the precleared super-

![Figure 6](https://iovs.arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/932963/)

**Figure 6.** Immuno-EM analysis of Als2cr4 in WT retinas. Retinas from either (A) light- or (B) dark-adapted mice were identified with pAb FLJ-FM and secondary goat anti-rabbit-10 nm colloidal gold antibody particles. (A, B) Rod OS. IM-EM shows Als2cr4 embedded within the discs of photoreceptor OS. Magnification, ×20,000.

![Figure 7](https://iovs.arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/932963/)

**Figure 7.** Membrane localization of Als2cr4 transfected into HEK293 cells. Confocal microscopy images of 6xHis tagged-Als2cr4, transiently transfected into HEK293 cells. Als2cr4 subcellular localization was identified by immunohistochemical labeling with antibodies recognizing Als2cr4 with pAb FLJ-FM (green; a, b) or the His-tag with mAb 6xHis (red; c, d). Nucleic acids in the nuclei were labeled with TO-PRO 3 iodide. Cells transfected with a recombinant Als2cr4 cDNA construct expressed the recombinant Als2cr4 protein, which is localized to the cytoplasm and the plasma membrane. (f) Whereas those transfected with no recombinant cDNA inserted into the pCDNA4/HisMax vector (e) exhibited no Als2cr4 expression, and the His-tag was limited to the nucleus. Labeling patterns of Als2cr4 pAb (b) and 6xHis mAb (d) displayed co-localization in the plasma membrane (f), thereby confirming Als2cr4 antibody specificity.
tant fraction, and lane 2 shows the proteins eluted after IP with pAb FLJ-FM (Supplementary Table S1, http://www.iovs.org/cgi/content/full/51/9/4407/DC1). Eight bands were excised and analyzed by mass spectra analysis (Table 1). Of the 15 peptides identified in the eight bands, most corresponded to cytoskeletal components. In contrast, the only bands detected in the control IP fraction were IgG light and heavy chains. Figure 8 shows IB analysis with specific antibodies verifying the direct interaction of Als2cr4 with myosin Va, myosin VI, and Arr4. Moreover, the observation that Als2cr4 immunoprecipitates Arr4 independently confirms the protein–protein interaction previously observed by the retinal yeast two-hybrid screen with cone arrestin used as the bait. Further experimental work is ongoing to determine the biological significance of this interaction. In addition, Als2cr4, the positive control, is clearly shown to be immunoprecipitated with FLJ-FM pAb and detected by the second Als2cr4 antibody FLJ-LG pAb; and Arr1, a negative control in this experiment, verifies that Arr1 binds nonspecifically to the protein G beads.

### DISCUSSION

Initially, visual Arr4 was used as a bait to screen a yeast two-hybrid Nrl−/− retinal cDNA library, and one of the candidates for protein–protein interaction was a novel hypothetical protein known as Als2cr4. Because of its high mRNA expression in the eye and the observation that Als2cr4 is originally a candidate gene for ALS, we explored the potential physiological relevance of this novel protein. In parallel to our work, researchers in a recent study tested the functionality of Als2cr4 and, in zebrafish knockdown studies with morpholino technology, demonstrated an “eyes absent” phenotype (The Zebrafish Model Organism Database, www.zfin.org).

Photoreceptor studies with confocal IHC and EM-EM revealed that Als2cr4 is targeted to the membrane and is concentrated in the rod OS. These data correlate well with our IB analysis (Fig. 2) showing lower Als2cr4 protein levels in retinas from “all cone” Nrl−/− mice. Moreover, Als2cr4 appeared to be embedded in the outer segment disc ring; however, whether it is present in the photoreceptor disc rim like peripherin/rds or in the flat lamellar disc region where opsin is localized is not known. These morphologic studies suggested that Als2cr4 contributes to either phototransduction protein trafficking or participates as a structural membrane protein, in agreement with previous proteomic observations.10,11 In ongoing experiments, we are examining Als2cr4 within the CC and the horizontal cells of the OPL.

Using computer modeling software, we predicted that Als2cr4 is related to the tetraspanin superfamily of proteins containing four transmembrane-spanning domains, two cytoplasmic tails, two extracellular loops, and one intracellular loop. However, Als2cr4 has major differences with canonical

---

**TABLE 1. Retinal Proteins Identified by Mass Spectrometry from Immunoprecipitation with Als4cr2 pAb FLJ-FM**

<table>
<thead>
<tr>
<th>Band No.</th>
<th>NCBI Accession No.</th>
<th>Protein ID</th>
<th>Mass (kDa)</th>
<th>Peptides (n)</th>
<th>Coverage (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2*</td>
<td>NP_780469</td>
<td>Myosin X</td>
<td>233</td>
<td>18</td>
<td>9</td>
</tr>
<tr>
<td>2*</td>
<td>NP_082297</td>
<td>Myosin XIV</td>
<td>229</td>
<td>10</td>
<td>6</td>
</tr>
<tr>
<td>3</td>
<td>NP_043994</td>
<td>Myosin Va</td>
<td>215</td>
<td>18</td>
<td>11</td>
</tr>
<tr>
<td>4</td>
<td>NP_001034635</td>
<td>Myosin VI</td>
<td>146</td>
<td>18</td>
<td>18</td>
</tr>
<tr>
<td>5</td>
<td>Q61768</td>
<td>Kinesin-1</td>
<td>110</td>
<td>7</td>
<td>9</td>
</tr>
<tr>
<td>6</td>
<td>NP_034851</td>
<td>Lamin-B1</td>
<td>67</td>
<td>5</td>
<td>9</td>
</tr>
<tr>
<td>7*</td>
<td>NP_035851</td>
<td>Vimentin</td>
<td>54</td>
<td>34</td>
<td>59</td>
</tr>
<tr>
<td>7*</td>
<td>NP_001032901</td>
<td>Als2cr4</td>
<td>45</td>
<td>6</td>
<td>16</td>
</tr>
<tr>
<td>8*</td>
<td>NP_031419</td>
<td>Beta-actin</td>
<td>42</td>
<td>16</td>
<td>47</td>
</tr>
</tbody>
</table>

Summary of the eight proteins isolated by immunoprecipitation with the FLJ-FM pAb and identified by mass spectrometry, their predicted molecular mass, the number of peptides identified for each protein, and the percentage of the protein covered.

* Bands that showed protein grouping ambiguity.
tetrascanspans: mainly, the presence of a shorter extracellular loop (29 AA vs. 100 AA) and the existence of a large amino-terminal tail (226 AA) compared with the relatively short amino-terminal tail of other tetrascanspans. In addition, tetrascanspans contain a characteristic four to six conserved cysteine residues in the large outer loop that Als2cr4 does not contain.  

In retinal disease, the most prominent members of the tetrascanspan family are peripherin/rds32–35 and ROM-1.34 These proteins participate in outer segment morphogenesis by providing a structural basis whereby new discs and lamellae are continually added to the photoreceptor outer segment basal region.32

The outer segment renewal is a critical process as photoreceptors undergo daily phagocytosis of apical discs and lamellae by retinal pigment epithelium. Moreover, forms of retinitis pigmentosa, macular degeneration, and macular dystrophy are linked to defective genes encoding peripherin/rds, ROM-1, and prominin (Prom1).35 In mouse models, Rds mutants fail to form rod OS,36 whereas in cones the phenotype is less severe with formation of enlarged and morphologically disorganized cone OS.24 In comparable studies, loss of ROM-1 leads to engorged discs and disordered rod OS.37 Furthermore, loss of Prom1 is associated with overgrown and misoriented disc membranes.38 As is evident from these studies, the loss of structural proteins has devastating effects on photoreceptor morphology and leads to loss of photoreceptors. Because Als2cr4 shares structural similarities, has a comparable immunostaining pattern,24,34 and is embedded in the discs, we hypothesize that, similar to peripherin/rds and ROM-1, this structural protein contributes to maintaining the structural integrity of photoreceptors.

To further decipher the protein biochemistry for Als2cr4, we looked at its interacting partners in the DA Nfr1/−/− retina with co-IP experiments with pAb FLI-FM (Supplementary Fig. S4, Supplementary Table S1). In total, eight bands were excised, subjected to mass spectra analysis, and identified: vimentin, actin, myosin Va, myosin VI, myosin X, myosin XIV, kinesin I, Als2cr4, and lamin B-1 (Table 1). The cytoskeleton has been implicated in many aspects of vision ranging from structural maintenance of photoreceptors to protein trafficking between the inner segment (IS) and OS and for active transport in the connecting cilium.39 Cilia are hairlike protrusions extending from the surface of cells involved in cellular motility or, in the case of primary cilia, specialized functions including sight, smell, and hearing. Ciliopathy is a disease state in which mutations are located in protein coding regions and are involved in cilia formation or protein transport. Primary ciliary dyskinesia, kidney diseases, respiratory diseases, and retinal degenerations are all examples of ciliopathies. In addition, mounting evidence links mutations in genes responsible for photoreceptor morphogenesis and intracellular transport (IFT) with photoreceptor degeneration.40–46 These diseases are usually accompanied by other clinical symptoms due to global detrimental effects on primary cilia and are syndromic.39

Like all ciliated cells, photoreceptor OS lack the organelles for protein synthesis and are thus dependent on protein trafficking. To date, there is an ongoing debate as to the major mode of shuttling signaling proteins between the inner and outer segment subcellular compartments. Some experts argue for diffusion,47 whereas others support active transport mediated by molecular motors along actin filaments or microtubules.48 Because of the abundance of cytoskeletal components interacting with Als2cr4 (Table 1, Fig. 8), and in particular molecular motors, we propose that Als2cr4 is actively transported to its destination by members of the kinesin and/or myosin families.

Investigation as to the etiology of Usher 1B syndrome,49 which is the most frequent disease causing combined deafness and blindness in humans, identified defects in the myosin VII gene. The absence of the molecular motor myosin VII is deleterious, as it plays a role in transporting rhodopsin and other proteins essential for normal phototransduction. The cytoskeleton plays an essential role in the photoreceptor’s maintenance by providing structure in the form of actin filaments and microtubules forming the axoneme. A further example is the loss of the protein reinitis pigmentosa 1 (RP1) that binds to and stabilizes microtubules and is involved in the regulation of axonemal length and stability.40 The cytoskeleton also plays an indispensable role in transport mechanisms, as is evident by the conditional knockout of kinesin (Kif3a), which results in the mislocalization of opsins, Arr1, and peripherin to the inner segment,50 and the IFT88 knockout mouse,51 which exhibits uncharacteristic stacking of disc membranes reminiscent of ROM-1 loss in rods or peripherin/rds loss in cones. In addition to irregular morphology, loss of IFT88 also leads to an accumulation of opsins in the outer segment and, ultimately, to cytotoxicity.

In conclusion, we have for the first time, characterized the novel protein Als2cr4 in the retina and proposed that it has a 2D structure consisting of four transmembrane domains. We also discovered this structural protein is embedded in the photoreceptor outer segment discs and in the postsynaptic OPL in horizontal cell axons. Furthermore, we identified cytoskeletal interacting partners and hypothesize that Als2cr4 functions in membrane trafficking and contributes to maintaining photoreceptor structure. Although additional experiments are essential for resolving the question of the function of Als2cr4 in the mouse retina, the recent morpholino knockdown experiment using the Als2cr4 orthologue (XP_685343) to demonstrate a significant phenotype with “failure to develop the eye” (The Zebrafish Model Organism Database www.zfin.org, NCBI gene ID Q66IE4) suggests that our hypothesis is valid. Future experiments will determine whether Als2cr4 functions analogously to what is observed in zebrafish retina as an early embryonic component that is indispensable for normal mouse eye development and for maintaining the photoreceptor structural integrity.

Acknowledgments

The authors thank Mary D. Allen for generous and continuous support; Jeannie Chen for the Arr1/−/− mice; Anand Swaroop and Jose Luis Linares for providing Nfr1/−/− mice and Nfr1/−/− cDNA library for the yeast two-hybrid screen; and Bruce Brown, Xumei Zhu, and Jeffrey Raskin for their technical and scientific expertise; Barbara Nagel for her immunoelectron microscopy; India Jane Bradley for her help with immunohistochemistry and confocal microscopy; and members of the Mary D. Allen Laboratory, including Shun-Ping Huang, Rosanne Yetemian, Leng-Ying Chen, and Guey Shuang Wu.

References


