Identification and Light-Dependent Translocation of a Cone-Specific Antigen, Cone Arrestin, Recognized by Monoclonal Antibody 7G6

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PURPOSE. To elucidate the antigen recognized by monoclonal antibody (mAb) 7G6, a widely used cone-specific marker.

METHODS. 7G6 immunocytochemistry was performed on sections of human, primate, and bovine retina. The antigen was immunoprecipitated from human retinal lysates and purified with protein G. Edman degradation and liquid chromatography of tryptic peptides combined with tandem mass spectrometry (LC-MS/MS) identified the antigen.

RESULTS. Sequencing of peptides derived from the immunoprecipitated 7G6 antigen identified it as cone arrestin. The identity was confirmed by Western blot analysis with recombinant human cone arrestin and competition with the antibody in immunocytochemistry. Subcellular localization of cone arrestin in dark-adapted and bleached bovine retinas showed that cone arrestin accumulated in cone outer segments of light-adapted retina but was more concentrated in the inner segments of dark-adapted retina. By expression of truncated human cone arrestin mutants systematically deleting areas divergent from bovine and primate cone arrestins, the epitope of 7G6 was identified as a divergent loop exposed at the surface within the N-domain of cone arrestin.

CONCLUSIONS. Several independent methods established that the 7G6 antigen is cone arrestin. The 7G6 epitope is contained in a divergent loop, the sequence of which is conserved in bovine and primates but not other vertebrate species consistent with the specificity of the antibody. The light-dependent translocation of cone arrestin suggests a role in light–dark adaptation of cones. Because of the location of its gene on the X-chromosome, cone arrestin is a candidate gene for X-linked cone dystrophies. (Invest Ophthalmol Vis Sci. 2003;44:2858–2867) DOI:10.1167/iovs.03-0072

The cone-specific monoclonal antibody (mAb) 7G6 has been shown to recognize all cone subtypes in the adult primate retina and the labeling for it has been observed throughout the cytoplasm of the cones.1 The antibody 7G6 has been used as a universal primate cone marker in normal and degenerating retina in several subsequent publications.2-5 (A teaching set on normal and diseased human retinas assembled by Ann Milam displaying retina sections stained with mAb 7G6 is available at http://www.penneye.com/html/jeff_berger_memorial.html.) The 7G6 antigen was recently shown to colocalize in the cone cytoplasm with an unidentified p45 protein that appears to share an epitope with the transcription factor NRL.4 During early primate development, some heterogeneity in 7G6 labeling was observed. For example, a subset of blue cones was delayed in their acquisition of 7G6 immunoreactivity, suggesting a delayed expression of the antigen, whereas the onset of 7G6 immunoreactivity in red and green cones preceded both the expression of cone opsins and the formation of synaptic contacts in the inner plexiform layer.1

To investigate the nature of the antigen and its gene more closely, we first used Western blot analysis of human macula components to determine the mobility of the antigen in SDS-PAGE. We then purified the antigen, generated proteolytic peptides and sequenced them by both Edman degradation and liquid chromatography combined with mass spectrometry (LC-MS/MS). Sequence analysis of multiple fragments showed unambiguously that the 7G6 antigen is indistinguishable from cone arrestin (Carr, or X-arrestin). (Zhang H, Cuenca N, Church-Kopish J, Ivanova T, Frederick JM, MacLeish PR, Baehr W, ARVO abstract 1414, 2002). Arrestins are a large gene family involved in regulation of G-protein–coupled receptors and G-protein-linked cascades. The cone’s visual cascade is thought to function very similarly to that of rods, but is composed of components encoded by distinct genes. Most of these components have been cloned and characterized, including cone pigments,5 cone transducin α-subunit,6 cGMP PDE5b,7 and cyclic nucleotide gated channel α(CNG)3,6 and β subunit.9 Most regulatory components such as guanylate cyclase-activating protein (GCAP)-1,10 guanylate cyclase (GC)-1,11 RGS9-1,12 G-protein–coupled receptor kinase (GRK)-1,13 and recoverin14,15 are present in both rods and cones in mammals. Exceptions are the arrestins, which occur in cell-specific isoforms.

One hallmark of arrestin function is its relocalization in the rod cell in a light-dependent manner. In dark-adapted retina, arrestin is located in the inner segment, but redistributes rapidly to the outer segment on illumination.16,17 Recently, cone arrestin was shown to translocate in the mouse retina, depending on illumination.18 We found that bovine cone arrestin redistributes similarly in light- and dark-adapted bovine retina. In the light, cone arrestin accumulates in the cone outer segments, presumably interacting with bleached cone visual pigments and quenching the cone cascade. In contrast, in the dark, cone arrestin accumulates in the inner segment. The light-dependent translocation of cone arrestin depending on...
light history, similar to that of rod arrestin, suggests a novel mechanism of light-dark adaptation in cones.

**Materials and Methods**

**Immunoprecipitation and Peptide Sequencing**

Retinal extracts were prepared from human macular buttons obtained from the Lions Eye Bank at the University of Utah 4 to 8 hours after death. A crude extract of human maculae was mixed with 7G6 antibody and protein-G–coupled agarose (Santa Cruz Biotechnology, Santa Cruz, CA) in IP buffer (20 mM Tris [pH 7.5], 150 mM NaCl, and 0.1% Tween 20). The mixture was incubated 2 hours at room temperature on a rocking platform. The agarose beads were pelleted by brief centrifugation and washed three times with IP buffer. SDS (2X) gel loading buffer (45 μL) was added to the agarose beads followed by boiling for 2 minutes to elute the bound proteins completely. The proteins were fractionated by 10% SDS-PAGE gel and stained with Coomassie blue dye. Stained bands corresponding to the antigen were excised from SDS-PAGE gel and submerged into destaining solution containing 0.4 μg sequencing-grade trypsin (Promega Corp., Madison, WI). The proteolysis was stopped by adding 1/10 volume of 10% trifluoroacetic acid (TFA). The released peptides were extracted several times from the gel with 100 μL of 0.1% TFA and 60% acetonitrile followed by concentration (Speed-Vac; Thermo Savant) concentration. After incubation at 30°C for 1 hour, the liquid was removed, and the fresh destaining solution was added. The process was repeated until the gels were completely destained. The gels were then fully dehydrated in a concentrator (SpeedVac; Thermo Savant, Holbrook, NY). For in-gel digestion, the dried gels were rehydrated with 40 μL 0.2 M NH4HCO3 containing 30% acetonitrile. After incubation at 30°C for 1 hour, the liquid was removed, and the supernatant was saved, and the pellet was homogenized with a plastic pestle in 100 μL hypotonic buffer A (20 mM Tris, [pH 8.0], 1 mM EDTA, 1 mM dithiothreitol [DTT], and 100 μM phenylmethylsulfonyl fluoride [PMSF]). The tissue lysate was centrifuged at 600g for 5 minutes at 4°C. The supernatant was removed and saved, and the pellet was homogenized again in buffer A and recentrifuged as before. The supernatants from these two centrifugations were combined and subjected to ultracentrifugation at 250,000g for 1 hour at 4°C. One tenth of the supernatant and the entire pellet from ultracentrifugation were subjected to 10% SDS-PAGE gel electrophoresis, and the resolved polypeptides were transferred to a nitrocellulose filter (Bio-Rad, Hercules, CA). The filter was probed with mAb 7G6 (1:150) followed by horseradish peroxidase [HRP]-conjugated secondary antibody. The signal was visualized using phosphorescence (ECL system; NEN Life Science, Boston, MA) and detected by x-ray film.

**Western Blot Analysis**

Human maculae were isolated from freshly obtained retinas using a trephine (6 mm diameter). Two maculae were dropped into a microcentrifuge tube and homogenized with a plastic pestle in 100 μL hypotonic buffer A (20 mM Tris, [pH 8.0], 1 mM EDTA, 1 mM dithiothreitol [DTT], and 100 μM phenylmethylsulfonyl fluoride [PMSF]). The tissue lysate was centrifuged at 600g for 5 minutes at 4°C. The supernatant was removed and saved, and the pellet was homogenized again in buffer A and recentrifuged as before. The supernatants from these two centrifugations were combined and subjected to ultracentrifugation at 250,000g for 1 hour at 4°C. One tenth of the supernatant and the entire pellet from ultracentrifugation were subjected to 10% SDS-PAGE gel electrophoresis, and the resolved polypeptides were transferred to a nitrocellulose filter (Bio-Rad, Hercules, CA). The filter was probed with mAb 7G6 (1:150) followed by horseradish peroxidase [HRP]-conjugated secondary antibody. The signal was visualized using phosphorescence (ECL system; NEN Life Science, Boston, MA) and detected by x-ray film.

**Immunocytochemistry and Preabsorption Assay**

Monkey eyes were obtained from the Texas Primate Center (Alice, TX) and fixed as described previously. Adult human and bovine eyes were dissected to remove the anterior segment, and the eye cup was transferred to freshly prepared 4% parafomaldehyde in 0.1 M phosphate buffer (pH 7.4). After fixation for 2 hours at room temperature, the eye cup was immersed in 10% sucrose in 0.1 M phosphate (1 hour), 20% sucrose in 0.1 M phosphate (1 hour), and 30% sucrose in 0.1 M phosphate buffer (overnight). A small piece of retina-RPE was isolated with a razor blade, embedded in commercial medium (TBS: Triangle Biomedical Sciences, Durham, NC) and frozen. Sections (12-μm thick) were cut with a cryostat and mounted on gelatin-coated or charged slides (Superfrost/Plus, Fisher Scientific, Pittsburgh, PA). Bovine sections from light- and dark-adapted specimens were mounted pair-wise so that tissues from both were incubated in the same pool of antibody solution. This facilitated direct comparison of 7G6 immunolocalization correlating with light-dark adaptation.

For immunocytochemistry, the sections were blocked with 0.1 M phosphate buffer (pH 7.4), containing 1% BSA, 0.1% Triton X-100, and 10% normal goat serum, and probed with 7G6 antibody (1:100 or 1:200 dilution) followed by FITC-conjugated goat anti-mouse or TRITC-conjugated goat anti-mouse secondary antibody. For double labeling, the sections were incubated simultaneously with mAb 7G6 and polyclonal antibodies anti-calbindin (Swant, Bellinzona, Switzerland) overnight at 4°C. Sections were visualized by application of rhodamine (TRITC)-conjugated goat anti-mouse and FITC-conjugated goat anti-rabbit secondary antibodies. The immunolocalization was imaged using a confocal microscope (model LSM 510; Carl Zeiss Meditec, Thornwood, NY) set to an optical slit of 1 μm.

A preabsorption study was performed to verify 7G6 antibody specificity. Fresh human retina was obtained from a 4-month-old infant whose eye was enucleated because of retinoblastoma (biopsy tissue provided by Nick Mamalis, University of Utah). The nontumorous portion of the retina-RPE was dissected 2.5 hours after enucleation, fixed for 2 hours in 4% paraformaldehyde in 0.1 M phosphate buffer (pH 7.4), and cryoprotected in sucrose. Frozen sections were blocked for 1 hour with 10% normal goat serum before incubation with mAb 7G6. As a negative control, 7G6 (1:200) was preabsorbed for 6 hours with an excess amount of recombinant human cone arrestin (~15 μg rCarRHis/50 μL solution) before the solution was applied to the sections of retina. Immunoreactive sites were visualized by incubation 1 hour in FITC-conjugated goat anti-mouse IgG (1:300, cat. #115-095-146; Jackson Immunoresearch, West Grove, PA) as the secondary antibody and imaged as described for immunocytochemistry.

**Expression and Purification of Human His-Tagged Cone Arrestin**

The human His-tagged cone arrestin (hcArrHis) expression construct was kindly provided by Cheryl Craft (University of Southern California). For this construct, full-length human cone arrestin cDNA was cloned into frame in an expression vector (pTRC-His B, Promega). The plasmid was transformed into Escherichia coli ER2556 (New England Biolabs, Boston, MA), and a single colony was inoculated into 5 mL Luria-Bertani (LB) medium and grown at 37°C. The overnight culture was diluted into 100 mL of and continued growing to an OD600 of 0.6t0 0.7. Isopropyl-β-D-thiogalactopyranoside (IPTG) was added to the culture to a final concentration of 1 mM to induce the expression of recombinant protein. After 5 hours of induction at 37°C, the culture was harvested. The cells equal to 1.5 mL of culture were lysed with lysoyme, and a fraction of the lysate was analyzed by Western blot with 7G6 antibody. Recombinant hCarRHis was purified from 500 mL of culture with a HisTrap column (Pharmacia/Amersham, Piscataway, NJ) according to the manufacturer’s instructions. The purified protein was verified by SDS-PAGE gel and quantified using a protein assay kit (Cat. no. 500-0006; Bio-Rad).

**Mutagenesis**

To prepare human cone arrestin deletion mutants, mutagenesis was performed with a kit (QuickChange; Stratagene, LaJolla, CA). Five deletion mutants (see Fig. 4A) introducing stop codons after codons were introduced.
7G6 immunonegative (results not shown). The function of try, as described earlier. The eye was dissected in normal room light, and the retina-RPE was light with an intensity of approximately 500 lux (normal room light).

Metal can sealed with black tape and dissected in a darkroom under abattoir. One eye was dark adapted for 3 hours on ice in a light-tight box. The mutation was confirmed by DNA sequencing (Prism capillary sequencer; Applied Biosystems). The IPTG-induced expression, purification of the truncated proteins and Western blot were performed, as described earlier.

**Determination of Arrestin Gene Structures**

Contigs from human (AL357752), mouse (AC091784), rat (AC095218), and zebrafish (AL590151) were identified by screening the National Center for Biotechnology Information (NCBI) high-throughput gene sequences (htgs) using cone arrestin protein sequences and tblastin (NCBI database). Gene structures were determined with cDNA or protein sequences as templates. The novel zebrafish cArr gene structure was derived using 15 zebrafish cArr expressed sequence tags (ESTs)—AW826248, AW826514, BI033517, BI033512, BI036049, BF937662, BF937816, BI938932, BG305632, BG306321, BG306785, BG306798, BI076094, BB879022, and BB879162—covering the entire mRNA from the transcription start point to the poly(A) tail. Splice variations were examined in all genes using partial cArr-specific ESTs. Software for DNA analysis was Omiga 2.0 (Accelrys, Cambridge, UK; available at http://www.accelrys.com/dstudio/ds_gene/) and DNA Star (DNASTar, Inc., Madison, WI; http://www.dnastar.com), both available commercially.

**Light and Dark Adaptation of Bovine Eyes**

A pair of freshly enucleated bovine eyes was obtained from a local abattoir. One eye was dark adapted for 3 hours on ice in a light-tight metal can sealed with black tape and dissected in a darkroom under dim red light. The retina-RPE was fixed in the dark with 4% paraformaldehyde solution. The other eye, placed on ice, was kept at room light with an intensity of approximately 500 lux (normal room light). The eye was dissected in normal room light, and the retina-RPE was fixed in 4% paraformaldehyde and processed for immunocytochemistry as described earlier.

**RESULTS**

**Specificity of mAb 7G6**

We first explored the cell specificity of the mAb 7G6 in human, bovine, and primate retina (Fig. 1). Staining human frozen sections with 7G6 (Fig. 1B) reveals expression of the antigen exclusively in cones, apparently distributed throughout the cell including cone pedicles, somata, and inner and outer segments. Human retina incubated with a polyclonal anti-calbindin antibody (Fig. 1A) reveals labeling in the cytoplasm of the long-wavelength (LW) cone inner segments, the cell bodies, and the synaptic pedicles. Staining in the outer segments with this antibody is relatively weak. Note that short-wavelength (SW) cones are only very faintly labeled with anti-calbindin. The identity of these cones as SW cones was recently confirmed by double labeling with anti-blue opsin antibody. Anti-calbindin also labeled bipolar cells which are 7G6 immunonegative (results not shown). The function of calbindin, an EF hand containing small Ca$^{2+}$ binding protein with widespread distribution in neurons, is unknown, but it has been suggested that it transports intracellular Ca$^{2+}$ and protects against Ca$^{2+}$ mediated apoptosis. When human retina sections were double labeled with anti-calbindin and 7G6 (Fig. 1C), synaptic pedicles and inner segments of LW cones, but not SW cones, showed colocalization. Apart from human (Figs. 1B, 1C) and monkey (Fig. 1E) retinas, only bovine cones were found to be immunopositive with mAb 7G6 (Fig. 1D). Labelled cones of other vertebrate retinas (mouse, ground squirrel, salamander) with 7G6 was unsuccessful (results not shown), indicating that the 7G6 epitope is only present in primate and bovine antigens. The results are consistent with mAb 7G6 recognizing an antigen present nearly exclusively in the cytoplasm of primate and bovine cone photoreceptors. Uniform labeling with 7G6 and variation in double labeling with anti-calbindin suggests that the 7G6 antigen is present in all cone subtypes.

**Identification of the 7G6 Antigen**

To identify the 7G6 antigen in cones, we used Western blots with macular extracts from human retina (the macula is rich in cone photoreceptors). Western analysis revealed the presence of the 7G6 antigen exclusively in the hypotonic supernatant, and its absence in the membrane pellet (Fig. 2A). The apparent mobility in denaturing SDS gels is approximately 44 to 46 kDa, independent of denaturing the extract at high or low temperatures. To immunoprecipitate the 7G6 antigen, the hypotonic macular extract was incubated with mAb 7G6 and protein-G-coupled agarose. The immuno precipitate was analyzed by SDS-PAGE (Fig. 2B) revealing IgG (Fig. 2B, arrow 1) and the 7G6 antigen (Fig. 2B, arrow 2). We then determined proteolytic peptide sequences of the immunoprecipitated 7G6 antigen by two independent methods: Edman degradation and LC-MS/MS. Edman degradation of a single proteolytic polypeptide yielded a sequence identical with a short fragment of human cone arrestin (cArr) at positions 166-175 (Fig. 2C). Separation of a mixture of proteolytic fragments by liquid chromatography and subsequent mass spectrometric analysis of individual peaks allowed identification of fragments covering essentially the entire cArr polypeptide, except for the N-terminal region (Fig. 2C). The results show complete identity of the 7G6 antigen with human cArr, a polypeptide of 388 amino acid residues. The calculated molecular mass of human cArr is 43 kDa, consistent with the apparent mobility of the 7G6 antigen (44-48 kDa, Fig. 2A). In control experiments, we expressed His-tagged human cArr in bacteria and analyzed the recombinant protein by Western blot analysis (Fig. 2D). The results show that the recombinant protein has an apparent mobility of 50 kDa, slightly larger than the 7G6 antigen (the recombinant protein carries a His tag), and is immunopositive with mAb 7G6. When the mAb 7G6 is preabsorbed with recombinant cArr, immunohistochemical labeling of cones with 7G6 is completely abolished (Fig. 2E). These results established unambiguously that the 7G6 antigen is cone arrestin.

**Light-Dependent Redistribution of Cone Arrestin**

The identification of the 7G6 antigen as cone arrestin raised the interesting question of whether cone arrestins redistribute in cones, as has been observed for rod arrestin (s-antigen) and other phototransduction components (for review, see Ref. 23). In dark-adapted mammalian retina, arrestin is located in the inner segment, but moves rapidly to the outer segment on illumination. This apparent movement was controversial for many years, but recently was shown to occur unambiguously with rod To. Because dark-adapted primate retinas are very difficult to obtain, we performed a series of experiments with light- and dark-adapted bovine retinas, the only mammalian species besides primates with 7G6 immunoreactivity (Fig. 1D). In dark-adapted bovine retinas, cone arrestin distributes...
throughout the cytoplasm of cones, but accumulates predominantly in the myoid region of cone inner segments (Fig. 3A), the region where biosynthesis takes place. In the light, cone arrestin accumulates strongly in the outer segments (Fig. 3B), presumably interacting with bleached cone visual pigments and inactivating the cone cGMP cascade. Accumulation of these components in distinct compartments of photoreceptors suggests the presence of a novel mechanism of photoreceptor light- and dark-adaptation in both rod and cone photoreceptors.

Cone Arrestin Genes

To investigate potential links to human retina disease and the diversity of cone arrestins, we next analyzed vertebrate cone arrestin gene structures. The human and mouse cone arrestin genes have been cloned, and their exon–intron arrangement has been determined.25,26 These genes were shown to consist of 17 exons (one noncoding exon is located in the 5′ untranslated region [UTR]). We identified human (AC068231, AL357752) and mouse (AC091784) genomic contigs in GenBank that verified the published gene structures (Figs. 4A, 4B).

The human contig AL357752 containing the entire cArr gene localized the gene to chromosome X (q13.2-21.1) which significantly narrowed the published locus (Xcen-Cq22).25 In addition to human and mouse, we determined the gene structures of rat (Rattus norvegicus) and zebrafish (Danio rerio) cArr from large anonymous contigs (AC095218, AL590151) containing the entire cArr genes. As expected, the rat cArr gene structure (Fig. 4C), consisting of 17 exons, was nearly identical with that of mouse and human (Figs. 4A, 4B). In contrast, the zebrafish cArr gene (Fig. 4C) consisted of only 16 exons and encoded a 362 amino acid polypeptide with a significantly truncated C-terminal region. The zebrafish cArr gene features an exon 15 that is only 4 bp (one amino acid) in length, separated from exons 14 and 16 by a 302 and 91 bp introns, respectively. Like the mammalian cArr genes, the first exon is located in the 5′UTR. The predicted RNA and amino acid sequences were verified by 15 independent ESTs (10 of which are shown in Fig. 4D) that cover the entire 5′UTR, the coding sequence, and the 3′UTR. In contrast to complex alternative splicing observed in human and mouse cArr genes,25,26 the zebrafish ESTs deposited in GenBank show no significant
variation in splicing (apart from one EST with a short deletion in the 3'UTR). Phylogenetic analysis of the polypeptide sequences reveals that rod and cone arrestins fall into two distinct, but closely related subfamilies, as has been shown previously.27 The 7G6 epitope, however, is only present in primate and bovine cone arrestins.

**Functional and Divergent Domains in Cone Arrestins**

Vertebrate rod and cone arrestin amino acid sequences are highly conserved (Fig. 5) suggesting that the overall three-dimensional structure of cone arrestin is very similar to that of rod arrestin. The 0.28-nm crystal structure of rod arrestin showed a bipartite molecule consisting of N- and C-domains.28 The sequence similarity among rod arrestins is 51% to 83% (compared with bovine arrestin), among cone arrestins 52% to 84% (compared to bovine cone arrestins). Bovine arrestin is 47% to 55% similar to all cone arrestins shown in Figure 5. The arrestin signature region (positions 57-75 in human cArr) is essentially unaltered among all arrestins. Two N-linked glycosylation sites (Fig. 5, gray boxes) are conserved in the rod and cone arrestins, but N-glycosylation has not been established in arrestins (one report suggested that a small population of rod arrestin may be glycosylated). Among vertebrate cone arrestins, several regions are divergent, the C-terminal region (last 30 residues) and three relatively short regions of approximately 10 to 20 residues in the N-terminal half and the central part of the molecule (Fig. 5, bars 1–4). Region 3 in cone arrestin is analogous to a linker connecting two β sheets in the N- and C-domains of arrestin, respectively. In rod arrestin, the C-terminal region is important for locking arrestin into an inactive state.29 Sequence divergence in this region suggests flexibility in the length to exert this important function.

The amino acid sequences in Figure 5 are from the following sources: hum_cArr, accession no. 13633229 (National Institutes of Health); bov_cArr, accession no. D8534036; mus_cArr, accession no. NP_573468; rat cArr, derived from contig AC095218; xen_cArr, accession PS148327; zebArr3 (Danio rerio), derived from a cDNA sequence extracted from an anonymous genomic contig, accession no. AL590151 (Sanger...
FIGURE 3. Redistribution of cone arrestin in bovine cones, depending on the light history. (A) Dark-adapted bovine retinas. mAb 7G6 labeled the synaptic pedicles, the somata, and the inner and outer segments of the cones. In the dark, the bulk of cone arrestin was concentrated in the myoid region, the place of biosynthesis. (B) Fully bleached bovine retina. The bulk of cone arrestin was in the outer segments, where phototransduction occurs. The confocal images shown are representative of experiments performed with three different pairs of eyes in the fall of 2001 and the spring and summer of 2002.
Identification of the mAb 7G6 Epitope

To determine the epitope for 7G6 in human cone arrestin, we systematically deleted the divergent regions, starting at the C-terminal end, and assayed the truncation mutants for 7G6 immunoreactivity by Western blot analysis. Because human and bovine cArr C termini are nearly identical and distinct from the C termini of other cone arrestins, we suspected that the epitope for mAb 7G6 may be located there, particularly because this region carries a number of charged residues and is predicted to be antigenic by exposure on the surface of the molecule (see Fig. 2 in Ref. 28). However, C-terminal deletion mutants in which 23 and 35 C-terminal residues were removed (H90044 and H90045 in Fig. 5), were still immunoreactive (Fig. 6, bottom). We then deleted residues 188-388 (H90043 in Fig. 5), 169-388 (H90042), and 98-388 (H90041) in cone arrestin, and only deletion mutant H90041 was shown to be immunoo-negative with 7G6 (Fig. 6, top). Inspection of the sequence alignment narrowed the antigenic determinant for 7G6 in human cArr to a short region in the N-domain. This region is nearly identical in human and bovine cArr, but divergent in rodent and other vertebrate cone arrestin sequences. We conclude that the epitope recognized by mAb 7G6 is located in this divergent loop exposed to the surface of cone arrestin.

DISCUSSION

We have shown that a widely used immunocytochemical cone marker recognizes cone arrestin, a regulatory component of the cone cascade. Using deletion mutants of human cone arrestin, we have shown that the antigenic determinant is located in a divergent surface-exposed loop in the N-domain of cone arrestin, assuming that cone arrestin has a three-dimensional structure that is very similar to rod arrestin. The high sequence conservation between rod and cone arrestins (Fig. 5) largely justifies this conclusion. The proposed antigenic determinant is nearly identical in bovine and human cone arrestin, but is divergent in other cone arrestin sequences, consistent with the presence of the 7G6 epitope only in human, primate, and bovine retinas.

The function of arrestins consists of desensitizing a G-protein-coupled receptor (in photoreceptors, the rod and cone pigments). Similar to rod arrestin (for review, see Ref. 30), the function of cone arrestin is presumed to consist of interacting with phosphorylated cone visual pigments and downregulating or quenching the cone cascade, thus contributing to desensitization and light adaptation.31 Regeneration of the rod and cone visual pigments is thought to require phosphorylation by a rhodopsin kinase termed GRK132 or by both GRK1 and GRK7.33,34

Recombinant cone arrestins show little affinity for phosphorylated or nonphosphorylated rhodopsin.35,36 In vitro work with recombinant mouse cone arrestin suggests that it interacts with chicken cone pigments in a light- and phosphorylation-dependent manner.18 Based on crystal structure data of rod arrestin, it is assumed that the receptor’s C-terminal tail
containing the phosphorylation sites acts as a switch to induce a conformational change in arrestins that enables binding to the receptor. A similar model for arrestin/receptor interaction was proposed in solution on the basis of small angle x-ray scattering experiments. Rod arrestin’s binding to the receptor is sensitive to alternative splicing and missense muta-
tions. A bovine arrestin splice variant (p44) without the C-terminal peptide encoded by the last exon can bind to unphosphorylated rhodopsin. Arrestin mutant R175Q binds to activated phosphorylated and unphosphorylated rhodopsin, and Arr (R175E) binds to bleached rhodopsin from which the C-terminal phosphorylation sites have been removed. Other residues (D30, R175, D296, and D305S based on the bovine arrestin sequence) have been identified that enable binding to nonphosphorylated rhodopsin. The mutant arrestins are presumably switched into the active conformation in the absence of rhodopsin phosphorylation. All these residues are well conserved in cone arrestins (Fig. 5, asterisks), suggesting a similar functional role of cone arrestin in quenching the cone cascade. The uniform distribution of cone arrestin throughout the cell indicates that it diffuses freely in the cytoplasm. We cannot exclude that cone arrestin may have additional functions unrelated to quenching the cone cascade.

Malfunctioning in receptor phosphorylation (in which there is no quenching of the cascade) has been linked to cardiac disease in transgenic animal models and to congenital stationary night blindness in humans. Patients with naturally occurring mutations in rod arrestin have Oguchi disease, a form of inherited stationary night blindness. In arrestin-knockout mice, photoresponses are prolonged, consistent with the absence of quenching. In addition, it has been demonstrated that excessive light can cause rod cell death in pigmented rod arrestin-knockout mice. In human, the cArr gene is located on the long arm of the X chromosome near the centromere (Xcen-Xq22). Because cone arrestin’s presumed quenching capacity must be vital for proper cone functioning, its gene is considered a candidate gene for cone dystrophies or macular degenerations. No defects in the cone arrestin gene have so far been linked to human retinal dystrophies. In preliminary experiments, we screened three patients with X-linked cone disease for disease-causing mutations, but no mutation was identified (results not shown).

The movement of arrestins and transducin subunits in rods and cones is an interesting phenomenon representing a novel mechanism of light–dark adaptation. In light, arrestins accumulate in the outer segments to quench the phototransduction cascades. Rod transducin subunits, in contrast, move in the opposite direction to desensitize the cascade. The traffic from the inner to the outer segments must pass through a narrow interconnecting cilium in both rods and cones. In Kif3A conditional knockout mice, arrestin and rhodopsin transport, but not transducin transport, was shown to be dependent on kinesin II, a ubiquitous molecular motor present in the cilium. Unresolved questions include the mechanisms that trigger the relocalization of these components and exactly which motors and motor subunits are involved in translocation.

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References


