Connexin45-Containing Neuronal Gap Junctions in Rodent Retina Also Contain Connexin36 in Both Apposing Hemiplaques, Forming Bihomotypic Gap Junctions, with Scaffolding Contributed by Zonula Occludens-1

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Mammalian retinas contain abundant neuronal gap junctions, particularly in the inner plexiform layer (IPL), where the two principal neuronal connexin proteins are Cx36 and Cx45. Currently undetermined are coupling relationships between these connexins and whether both are expressed together or separately in a neuronal subtype-specific manner. Although Cx45-expressing neurons strongly couple with Cx36-expressing neurons, possibly via heterotypic gap junctions, Cx45 and Cx36 failed to form functional heterotypic channels in vitro. We now show that Cx36 and Cx45 coexpressed in HEK293 cells were colocalized in immunofluorescent puncta between contacting cells, demonstrating targeting/scaffolding competence for both connexins in vitro. However, Cx36 and Cx45 expressed separately did not form immunofluorescent puncta containing both connexins, supporting lack of heterotypic coupling competence. In IPL, 87% of Cx45-immunofluorescent puncta were colocalized with Cx36, supporting either widespread heterotypic coupling or bihomotypic coupling. Ultrastructurally, Cx45 was detected in 9% of IPL gap junction hemiplaques, 90–100% of which also contained Cx36, demonstrating connexin coexpression and cotargeting in virtually all IPL neurons that express Cx45. Moreover, double replicas revealed both connexins in separate domains mirrored on both sides of matched hemiplaques. With previous evidence that Cx36 interacts with PDZ1 domain of zonula occulden-2 (ZO-1), we show that Cx45 interacts with PDZ2 domain of ZO-1, and that Cx36, Cx45, and ZO-1 coimmunoprecipitate, suggesting that ZO-1 provides for coscaffolding of Cx45 with Cx36. These data document that in Cx45-expressing neurons of IPL, Cx45 is almost always accompanied by Cx36, forming “bihomotypic” gap junctions, with Cx45 structurally coupling to Cx45 and Cx36 coupling to Cx36.

Key words: double replica; FRIL; heterotypic coupling; homotypic coupling; PDZ domains; SDS-FRL

Introduction

In the retinal inner plexiform layer (IPL), rod and cone pathways are linked to output ganglion cells via abundant gap junctions...
Table 1. Antibodies used for Western blotting and immunohistochemistry

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<thead>
<tr>
<th>Antibody</th>
<th>Type</th>
<th>Species</th>
<th>Epitope; designation</th>
<th>Dilution (µg/ml)</th>
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<td>Whole protein; Ab06-332</td>
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<td>Millipore</td>
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</table>

aa, Amino acids.

- Antigen patterns used in situ hybridization and single-cell reverse transcription (RT)-PCR for connexin mRNA detection, or used the promoter sequences for Cx45 and Cx36 to drive expression of either lacZ (β-galactosidase) or enhanced green fluorescent protein (EGFP).
- The results appeared to demonstrate that many neurons express either Cx36 or Cx45, but coexpression of these connexins was not detected in any retinal neurons (Han and Massey, 2005; Schubert et al., 2005; Sohl et al., 2005; Dedek et al., 2006), except minimal in one study (Lin et al., 2005).
- Further, Han and Massey (2005) reported that “cx45 immunopuncta never colocalize with cx36 puncta” and suggested that Cx36 and Cx45 form separate populations of homotypic gap junctions. In contrast, Dedek et al. (2006) reported that 30% of Cx45-immunopositive puncta were colocalized with Cx45 and proposed those puncta to represent heterotypic Cx36-to-Cx45 gap junctions, which could account for abundant electrical and tracer coupling of neurons separately expressing Cx36 versus Cx45 (Maxeiner et al., 2005). That proposal raised difficulties because dye-coupling evidence suggested that Cx36 and Cx45 are unable to form functional heterotypic channels in HeLa cells (Teubner et al., 2000), which led to the suggestion that HeLa cells may lack essential coupling factors that otherwise permit heterotypic coupling of these connexins between IPL neurons (Dedek et al., 2006).
- The above discrepancies led us to investigate neuronal gap junctions in rodent IPL according to connexin composition, ultrastructural configurations, and complement of gap junction scaffolding and regulatory proteins ZO-1, ZO-2, and ZONAB. We then analyzed Cx36/Cx45/ZO-1 cellular localization and molecular interactions in HeLa expression systems and in mouse retina using immunoprecipitation (co-IP) and pull-down assays. Finally, we used freeze-fracture replica immunogold labeling (FRIL) (Rash et al., 2001) and SDS-digested freeze-fracture replica labeling (SDS-FRL) (Fujimoto, 1995) to colocalize Cx45 and Cx36 in hemiplaques of ultrastructurally defined gap junctions, including in double replicas of the same gap junctions. Our results reveal that most if not all Cx45-containing gap junction hemiplaques also contain Cx36, forming Cx36-to-Cx36 plus Cx45-to-Cx45 bihomotypic gap junctions.

Materials and Methods

- Antibodies and animals. Antibodies, their sources, and conditions under which they were used in this study are presented in Table 1. Most antibodies were obtained from Invitrogen, including a recently available polyclonal antibody generated against a peptide corresponding to a sequence within the C terminus region of mouse Cx45. Mouse monoclonal anti-Cx45 was also obtained from Millipore Bioscience Research Reagents. Specificities of the anti-Cx36, anti-ZO-1, anti-ZO-2, and anti-ZONAB antibodies have been demonstrated previously (Li et al., 2004a; Penes et al., 2005; Ciolofan et al., 2006). Rabbit anti-glutathione S-transferase (GST) Ab06-332 for detection of GST-PDZ domain fusion proteins was obtained from Millipore. Monoclonal and polyclonal anti-FLAG antibodies used as control IgGs in communoprecipitation studies were obtained from Sigma-Aldrich. For immunocytochemical studies, we used 35 adult male CD1 mice, five male C57BL/129SvEv wild-type and five Cx45 knock-out (ko) mice. For FRIL, we used five adult male and five adult female mice [four albino mice expressing GFP in locus ceruleus neurons (van den Pol et al., 2002; Rash et al., 2007b) (mice courtesy of Dr. Anthony van den Pol, Yale University, New Haven, CT), five C57BL/6j, and one C3L3], plus three adult male Sprague Dawley rats. One Wistar rat was used for SDS-FRL double replicas. All experiments were conducted according to protocols approved by our respective institutional Animal Care Committees, as prescribed in the Principles of Laboratory Animal Care (National Institutes of Health publication No. 86-23, revised 1985), with minimization of stress to animals and minimization of the number of animals used.

- Light microscope immunohistochemistry. Mice deeply anesthetized with equithesin (3 ml/kg) under photopic conditions were transcardially perfused with 3 ml of cold 50 mM sodium phosphate buffer, pH 7.4, containing 0.9% NaCl, 0.1 M sodium nitrite, and 1 U/ml heparin, and then with 40 ml of cold 0.16 M sodium phosphate buffer, pH 7.6, containing 1% formaldehyde and 0.2% picric acid. To ascertain the extent to which the total exposure to formaldehyde alters detectability of connexins, additional mice were perfused with half as much fixative for half the time. In both cases, residual fixative was immediately flushed from animals by perfusion with 10 ml of 25 mM sodium phosphate buffer, pH 7.4, containing 10% sucrose, thereby terminating further formaldehyde exposure. By comparison, other laboratories routinely fix via perfusion with 4% formaldehyde for 10 min (Maxeiner et al., 2005), immersion of Vibratome slices of retina in 4% formaldehyde for 10 min (Güldenagel et al., 2000), or immersion of isolated retinas in 4% formaldehyde for 30 min (Lin et al., 2005). These differences may be important because some investigators detected no Cx45 immunofluorescence in the OFF sublamina of IPL (Güldenagel et al., 2000), whereas other groups detected Cx45 in OFF sublamina and greater density of puncta in ON sublamina (Han and Massey, 2005).

- After perfusion fixation, eyes were removed, cryoprotected for 24–72 h in the final rinse, supplemented with sodium azide. Vertical sections of retina were cut (10 µm) on a cryostat and collected on gelatinized glass slides. Sections were washed for 20 min in 50 ml Tris- HCl, pH 7.4, containing 1.5% sodium chloride (TBS) and 0.3% Triton X-100 (TBST). For double-immunofluorescence labeling, sections were incubated simultaneously with two primary antibodies for 24 h at 4°C, then washed for 1 h in TBST, and incubated for 1.5 h at room temperature simultaneously with appropriate combinations of secondary antibodies, which included the following: FITC-conjugated horse anti-mouse IgG diluted 1:100 (Vector Laboratories), Cy3-conjugated goat anti-mouse IgG di-
luted 1:200 (Jackson ImmunoResearch Laboratories), Alexa Fluor 488-conjugated goat anti-rabbit IgG diluted 1:1000 (Invitrogen), and Cy3-conjugated donkey anti-rabbit diluted 1:200 (Jackson ImmunoResearch Laboratories). All antibodies were diluted in TBS Tr containing 5% normal goat or normal donkey serum. After secondary antibody incubations, sections were washed in TBS Tr for 20 min; in 50 mM Tris-HCl buffer, pH 7.4, for 30 min; covered with antifade medium; and coverslipped. Some sections single labeled with anti-Cx45 or anti-Cx36 were counterstained with either red Nissl fluorescent NeuroTrace (stain N21482) or green Nissl NeuroTrace (stain N21480) (Invitrogen). Control procedures included omission of one of the primary antibodies with inclusion of each of the secondary antibodies to establish absence of inappropriate cross-reactions between primary and secondary antibodies or between different combinations of secondary antibodies.

For immunolabeling of HeLa cells grown on poly-t-lysine-treated glass coverslips, cells were fixed with ice-cold 1% or 2% formaldehyde for 5 min, washed in TBS, and incubated for 16 h at 4°C simultaneously with rabbit anti-Cx45 and mouse anti-ZO-1, with mouse anti-Cx45 and rabbit anti-Cx36, or with mouse anti-Cx45 and rabbit anti-Cx36 in TBS containing normal donkey serum. Cultures were then washed in TBS Tr for 1 h and incubated for 1 h at room temperature simultaneously with FITC-conjugated donkey anti-rabbit IgG diluted 1:200 and Cy3-conjugated goat anti-mouse IgG diluted 1:400, or alternatively, with FITC-conjugated horse anti-mouse IgG diluted 1:200 and Cy3-conjugated donkey anti-rabbit IgG diluted 1:400. Slides were washed for 1 h in TBS Tr, covered with antifade medium, and coverslipped.

Conventional and confocal immunofluorescence images were acquired on a Zeiss Axioskop2 fluorescence microscope using Axiosvision 3.0 software and on an Olympus Fluoview IX70 confocal microscope using Olympus Fluoview 2.1 software, and assembled using Adobe Photoshop CS (Adobe Systems), CorelDRAW Graphics Suite 12 (Corel), and Northern Eclipse software (Empix Imaging). Some confocal images are presented as z-stacks of five confocal scans at z scanning intervals of 0.4 µm. However, quantitative analyses of colocalization relationships involving counts of puncta immunopositive for Cx45, Cx36, ZO-1, ZO-2, and ZONAB in vertical sections of the IPL were conducted using single confocal scans with a 60× oil-immersion objective lens and a digital zoom magnification of 4× or 5×. Counts were obtained from 4–10 separate, randomly selected fields along a lateral span of 55 µm encompassing the inner to outer edge of the IPL in retinas from two to four mice. Values for percentage colocalization were determined for each field and expressed as mean ± SEM. Numbers of Cx45-positive puncta in the IPL of wild-type and Cx36 ko mice were also determined and evaluated for statistically significant differences using unpaired Student’s t test.

For quantitative analysis of the effects of fixation strength and duration, immunofluorescence preparation was used that “clumped” collected from mice perfused with 40 ml of the above described 1% formaldehyde/0.2% picric acid fixative and compared with that obtained from mice perfused with 20 ml of the same fixative, with residual fixative immediately flushed in both cases. Double labeling was conducted using monoclonal anti-Cx45 (Millipore Bioscience Research Reagents) in combination with either polyclonal anti-Cx36 51-6300 or 36–4600 (Invitrogen), or alternatively, using polyclonal anti-Cx45 (Invitrogen) with monoclonal anti-Cx36 37-4600 (Invitrogen). As data obtained with these alternative antibody combinations were not significantly different, data were pooled. Because many Cx45-positive puncta (labeled green) in images of IPL were very small, overlay of these images with those of Cx36 (labeled red) were, in cases of Cx45/Cx36 colocalization, often insufficient to produce readily discernible yellow puncta that result from spatial overlap of green and red pixels, hence reducing confidence of establishing degree of colocalization by counting yellow puncta in superimposed images. Instead, quantitative data were acquired by transferring the locations of green Cx45-puncta onto transparencies and overlaying these transparencies onto images of red puncta representing the more abundant labeling for Cx36.

As a further test for Cx36/Cx45 colocalization in the retina versus simply coincidental overlap of these connexins resulting from the very high density of Cx36 in the IPL, images of Cx36 from the same field double labeled for Cx45 and Cx36 were flipped horizontally (Han and Massey, 2005), and Cx45-puncta overlapping with Cx36-puncta were again counted.

FRIL and SDS-FRL. SDS-FRL (Fujimoto, 1995, 1997) and FRIL (originally named by Grujters et al., 1987; as modified by Rash and Yasumura, 1999 and Rash et al., 2001) provide for high-resolution immunocytochemical identification of membrane proteins and simultaneous ultrastructural visualization of those proteins in distinctive membrane arrays. SDS detergent, as used in both FRIL and SDS-FRL, does not wash away many cytoplasmic scaffolding proteins that are strongly bonded to target transmembrane proteins in platinum/carbon-coated replicas, allowing cadherins, catenins, ZO-1, ZO-2, ZONAB, and PSD-95 to be immunogold labeled and localized to tight junctions, gap junctions, adherens junctions, and postsynaptic densities (Fujimoto, 1995; Fujimoto et al., 1997; Giolofan et al., 2006; Kamasawa et al., 2006). We used SDS-FRL for preparation of matched double replicas (see below) and FRIL for preparation of noncomplementary replicas. FRIL combines SDS-FRL with Lexan stabilization (Steere and Erbe, 1983) and “confocal grid-mapped freeze fracture” (Rash et al., 1995, 1997) for precise histological and ultrastructural mapping of immunogold-labeled gap junctions over square millimeter areas of complex CNS tissues.

For FRIL experiments, both mice and rats were used because previous immunofluorescence studies compared data from both species and reported minor but potentially significant species-specific differences (Feigenspan et al., 2001). In addition, we have published a large FRIL database characterizing distributions and morphologies of >2100 Cx36-containing gap junctions in the IPL of rats and >600 Cx36-containing gap junctions in IPL of mice, establishing the value of both species for comparative ultrastructural and immunocytochemical analyses under a variety of illumination conditions (Kamasawa et al., 2006). Mice and rats were deeply anesthetized with ketamine/xylazine and fixed by transcardiac perfusion with 2% or 4% formaldehyde, as described previously (Kamasawa et al., 2006). After dissection and collection of Vibratome sections cut at 150 µm, retina specimens were cryoprotected with 30% glycerol, frozen, freeze-fracture replicated, washed with SDS detergent (Fujimoto, 1995), and immunogold labeled, as described previously (Rash et al., 2001; Kamasawa et al., 2006).

For FRIL, the same antibodies used for light microscopy (above) were diluted to 10 µg/ml and applied for 1.2–3 h at either 8 or 22°C, and secondary antibodies were diluted 1:100 and applied for 11–15 h at either 8 or 22°C. We found that this level of dilution, along with an improved “labeling blocking buffer” (Dinckuh et al., 1987) significantly decreases “background,” with only minimal decrease in labeling efficiency (LE), yielding signal-to-noise ratios >15,000:1 (Meier et al., 2004). The resulting relatively low labeling efficiency (LE = 1:10 to 1:30, or one gold bead per 10–30 connexons) has the dual advantage that gold labels are sufficient to distinguish gold signal in the gold-labeled electron micrographs, and the gaps corresponding to gap junctions are clearly visible because of the high contrast of the label over the background. These features facilitate identification of the identity of the protein involved in the gap junction, as well as the selection of small areas of the retina where there is no or minimal background labeling (Rash et al., 2007).

To increase immunogold labeling efficiency, thereby potentially increasing detection of one or more connexins in “miniature” gap junctions (i.e., those containing <100 connexons) (Rash et al., 2007b), particularly those containing both Cx36 and Cx45, a second set of replicas (see Table 3) was labeled for 4.4–5.3 h in primary (vs our normal 1.2–3 h) and for 22 h in secondary (vs 11–15 h) antibodies. By way of comparison, previous light microscopic immunocytochemical studies labeled samples with primary antibodies overnight (>10 h) at 4°C (Dedek et al., 2006) or for as long as 3–5 d at 4°C (Han and Massey, 2005; Lin et al., 2005), and for 2–3 h at room temperature (Lin et al., 2005) to overnight at 4°C for secondary antibodies (Han and Massey, 2005; Dedek et al., 2006). Thus, FRIL experiments were at the lower concentration ranges and labeling times normally used for light microscopic and ultrastructural
electrophoresis in a 1.5% agarose gel prepared using TAE buffer (40 mM Tris-HCl, 0.1% acetic acid, 1 mM EDTA, pH 8.0, and 0.0004% ethidium bromide). Separated PCR products were isolated and excised under UV illumination and purified using a gel purification kit (Qiagen). PCR products were subcloned into PCR 2.1 vector, digested with BstXI, and ligated into pcDNA3 expression vector (Invitrogen) using T4 DNA ligase (Invitrogen) according to the manufacturer’s instructions. Cx45 recombinant plasmids were extracted and digested with XhoI to confirm insert orientation and then transfected with M13 forward and reverse primers for sequence confirmation.

Cell culture and HeLa cell transfection. HeLa cells (American Type Culture Collection), grown in DMEM supplemented with 10% fetal bovine serum and 1% penicillin-streptomycin, were transiently transfected with pcDNA3 vector or Cx45-pcDNA3 plasmids using LipofectAMINE 2000 reagent (Invitrogen), as described previously (Li et al., 2004a,b,c). HeLa cells were similarly cotransfected with Cx45 and either full-length Cx36 or truncated Cx36 that lacks the 4 C-terminal amino acids representing the PDZ domain-binding motif of Cx36 (prepared as described previously (Li et al., 2004a)). Transiently transfected HeLa cells were taken for analysis after a 24 h transfection period. For selection of clones stably transfected with Cx45, transfected cells were grown for 24 h and then passaged at 1:40 dilution with fresh DMEM containing 1.5 mg/ml G418, and individual clones expressing Cx45 were isolated after 3 weeks in culture. HeLa cells stably expressing Cx36 were prepared as described previously (Li et al., 2004a). Cocultures of HeLa cells stably expressing Cx36, Cx43, or Cx45 (generated as above or provided to us by Dr. K. Willecke, University of Bonn, Bonn, Germany) were prepared by mixing equal numbers of Cx36- and Cx45-expressing cells at a density of ~10,000 cells per chamber (1–1.5 cm²) on culture slides for 2 d or until reaching confluence. Transient transfection of Cx45 into HeLa cells stably expressing Cx36 was conducted as described above for transfection of HeLa cells with Cx45.

Western blotting. Deeply anesthetized animals were decapitated, and retinas were dissected, rapidly frozen, and stored at ~80°C until use. Cx45-transfected and vector-control HeLa cells were rinsed with PBS buffer (50 mM sodium phosphate buffer, pH 7.4, 0.9% saline), and these cells and retinas were homogenized in IP buffer (20 mM Tris-HCl, pH 8.0, 140 mM NaCl, 1% Triton X-100, 10% glycerol, 1 mM EDTA, 1.5 mM MgCl₂, 1 mM diithiothreitol, 1 mM phenylmethylsulfonyl fluoride, and 5 μg/ml each of leupeptin, pepstatin A, and aprotinin) and sonicated. Homogenates were centrifuged at 20,000 × g for 5 min, and the supernatants were taken for protein determination using a kit (Bio-Rad Laboratories). Western blotting was performed as described previously (Li et al., 2004a,b; Chen et al., 2006). Proteins (50–100 μg of protein per lane from tissue homogenates and 3 μg per lane from culture cell lysates) were separated by SDS-PAGE using either 10% or 12.5% gels followed by numbers of Cx36- and Cx45-expressing cells at a density of ~10,000 cells per chamber (1–1.5 cm²) on culture slides for 2 d or until reaching confluence. Transient transfection of Cx45 into HeLa cells stably expressing Cx36 was conducted as described above for transfection of HeLa cells with Cx45.

IP. Mouse retinas and HeLa cell lysates were homogenized in IP buffer, sonicated, and centrifuged at 20,000 × g for 10 min at 4°C. After protein determination of sample supernatants, volumes containing 1 mg of protein from mouse retina or from connexin-expressing HeLa cells were preclaried for 1 h at 4°C using 20 μl of protein-A-coated agarose beads (Santa Cruz Biotechnology), centrifuged at 20,000 × g for 10 min at 4°C, and incubated with 2 μg of polyclonal anti-ZO-1 antibody, 2 μg of monoclonal anti-Cx45 antibody, or 2 μg of polyclonal anti-Cx36 antibody for 1 h at 4°C. The mixture was incubated for 1 h at 4°C with 20 μl of protein-A-coated agarose beads and centrifuged at 20,000 × g for 10 min, and the pellet was washed five times with 1 ml of wash buffer (20 mM Tris-HCl, pH 8.0, 150 mM NaCl, and 0.5% NP-40). Samples were then mixed with an equal
volume of SDS-PAGE loading buffer (125 mM Tris-HCl, pH 6.8, 20% glycerol, 0.3 mM bromophenol blue, 0.14 M SDS, and 20% β-mercaptoethanol). For detection of Cx45, samples were boiled for 5 min and taken for immunoblotting with monoclonal anti-Cx45 antibody. For detection of Cx36, samples (not boiled) were immunoblotted with polyclonal anti-Cx36 antibody. Control samples were precipitated with omission of anti-ZO-1, anti-Cx36, or anti-Cx45 antibodies.

**Results**

**Distribution of Cx45 and Cx36 in retina**

Low-magnification images of vertical retinal sections labeled by immunofluorescence and counterstained with Nissl fluorescent NeuroTrace indicate the distributions and relative densities of Cx45 and Cx36 in mouse retina, where immunolabeling consisted exclusively of fluorescent puncta (Fig. 1). Cx45 immunofluorescence was faintly detectable in the outer plexiform layer (OPL) (Fig. 1), whereas Cx36-immunofluorescent puncta were substantially more abundant. Both monoclonal anti-Cx45 antibodies from Invitrogen and Millipore Bioscience Research Reagents, as well as polyclonal anti-Cx45 antibody (Fig. 1), produced punctate labeling throughout the IPL. The polyclonal antibody produced additional sparse but robust labeling at the outer boundary of the inner nuclear layer (INL) (Fig. 1C), as well as punctate labeling along blood vessels in retina (Fig. 1D) and brain (Fig. 1E), consistent with previous descriptions of Cx45 distribution in rodent retina (Dedek et al., 2006) and in cerebral vasculature smooth muscle (Krüger et al., 2000; Li and Simard, 2001). The additional labeling detected with the polyclonal antibody at known tissue locations of Cx45 expression (e.g., blood vessels) suggests that the epitopes of Cx45 on blood vessels are not detected by the anti-Cx45 monoclonal antibodies. Labeling with each of the anti-Cx45 antibodies was observed using optimized fixative containing 1% formaldehyde plus 0.2% picric acid. Greater formaldehyde concentration or greater fixative volume and longer duration of fixation, each followed by washout of residual fixative, substantially suppressed both labeling density and intensity, particularly for very small Cx45-positive puncta, as determined...
by the lower density of punctate immunolabeling of Cx45 in mice perfused with 40 ml compared with 20 ml of fixative (described below).

In the IPL, Cx36-positive puncta were consistently present at a higher density than Cx45-positive puncta, with the former outnumbering the latter by a ratio of \(\sim 2:1\). Using anti-Cx45 MAB3101 in combination with polyclonal anti-Cx36 antibodies in mice perfused with 20 ml of fixative, counts of puncta yielded \(736 \pm 27\) Cx36-puncta per field and \(427 \pm 32\) Cx45-puncta per field along a \(55 \times 55 \mu m\) \(x\)- and \(y\)-axis span of the IPL, with the latter encompassing the full vertical dimensions of the IPL. Labeling for Cx45 compared with Cx36 appears much less than reflected by these numbers because of the much smaller size of Cx45-puncta and the presence of what appeared to be aggregates of Cx36-puncta that could not be separately resolved because of the close proximity of gap junctions in the IPL (Kolb, 1979). This almost certainly led to a vast underestimate of the number of gap junctions containing only Cx36 (Kamasawa et al., 2006), and here, to underestimates of the ratio of Cx36- to Cx45-puncta. Cx45-puncta were in greater abundance in the inner compared with the outer part of the IPL (Fig. 1A). According to criteria for assigning dimensions of the IPL to inner and outer sublamina (Ghosh et al., 2004) and from counts of Cx36-puncta in confocal images obtained from retinas of mice, the inner three-fifths of the IPL, designated the ON sublamina, contained an average of \(297 \pm 13\) Cx45-puncta along a \(55 \mu m\) \(x\)- and \(y\)-axis span of IPL, whereas the outer two-fifths, designated the OFF sublamina, contained an average of \(130 \pm 31\) Cx45 puncta along the same span. Thus, the OFF sublamina has \(\sim 66\%\) of the volume density of Cx45 puncta as in the ON sublamina. Similar patterns of labeling for Cx45 in the IPL were obtained with polyclonal anti-Cx45. Labeling for Cx36 was densely distributed in the IPL, where it was more concentrated in the inner than the outer half, and moderately concentrated in the OPL (Fig. 1B). By way of comparison, the density of Cx45 puncta in the OFF sublamina in the briefly fixed samples was higher than in previously published studies, one of which reported an absence of Cx45 in the OFF sublamina (Gu¨ldenagel et al., 2000). Specifically, as reported by Dedek et al. (2006) using single confocal scans from mouse retina and using similar objective lenses as used by us, the density of Cx45-puncta in the OFF and ON sublamina of the IPL was \(0.0128\) and \(0.0242\) per \(\mu m^2\), respectively, whereas we found \(0.043\) and \(0.099\) per \(\mu m^2\), respectively. The density of Cx36-puncta they reported in the OFF and ON sublamina of the IPL was \(0.0168\) and \(0.0554\) per \(\mu m^2\), respectively, whereas we found \(0.097\) and \(0.146\) per \(\mu m^2\), respectively. These data indicate 3.4-fold to 4.1-fold difference in
detection of immunofluorescent puncta for Cx45 in OFF versus ON layers of IPL and a 5.8-fold to 2.6-fold difference in detection of Cx36. This greater detection of Cx36 cannot be attributable to nonspecific immunolabeling because all immunoreactivity with anti-Cx36 antibodies we used was absent in retina of Cx36 ko mice (described below).

**Immunofluorescence colocalization of Cx45 and Cx36 in the IPL**

Confocal double-immunofluorescence analysis was undertaken to compare the localization of Cx45 with Cx36 in the IPL (Fig. 2). In general, most puncta immunopositive for the two connexins in the ON sublamina were larger than those in the OFF sublamina, and there appeared to be a decrease in size and density of puncta from the...
ON to the OFF sublamina (Fig. 2A1, A2). Retinas from 12 mice were examined, and similar results were obtained from each. Initial quantitative confocal double-immunofluorescence analyses of Cx45 colocalization with Cx36 were conducted using retinal sections from animals transcardially perfused with 40 ml of fixative. With this fixation protocol, 46% of Cx45-puncta were found to be colocalized with Cx36-puncta (Table 2), approximating the values reported by Dedek et al. (2006). Because this value was substantially lower than the extent of Cx45 colocalization with Cx36 observed by FRIL (described below), we considered the possibility that some labeling of Cx45 was suppressed even with the already weak tissue fixation used (data not shown). Thus, quantitative analysis was repeated using retinas from mice perfused with 20 ml of tissue fixative, resulting in a proportional reduction in fixation time because residual fixative in all cases was flushed from animals by perfusion with buffered su- crose. This fixation protocol gave substantially greater numbers of Cx45-puncta and could not be separately resolved by confocal microscopy. This fixation protocol gave substantially greater numbers of Cx45-puncta and could not be separately resolved by confocal microscopy. In contrast to the results of Dedek et al. (2006), who found only minor overlap of Cx45 with Cx36 in the OFF sublamina of IPL, our image overlays revealed colocalization of Cx45 with Cx36 in virtually all regions of IPL, and showed that 87% of Cx45-puncta overlapped with Cx36-puncta (Fig. 2A, Table 2). The vast majority of these puncta showed total overlap, rather than partial overlap of adjacent puncta. Conversely, given the greater density of Cx36 than Cx45 in the IPL, most Cx36-puncta were not labeled for Cx45 (not quantified).

Partially overlapping or adjacent puncta were suggested as evidence for heterotypic gap junctions, with Cx36 on one side and Cx45 on the other side (Dedek et al., 2006). However, immuno- fluorescence labeling for Cx45 and Cx36 in apoposing hemi- plaques of truly heterotypic gap junctions would precisely over- lap and could not be separately resolved by confocal microscopy because of the molecular dimensions of apposed connexons versus the ~0.2 μm limit of resolution of light microscopy. To test whether published images showing “adjacent” puncta for Cx36 and Cx45 are consistent with close association of separate Cx36-homotypic and Cx45+Cx36 bimodulatory gap junctions at triadic appositions of cone bipolar cells with All amacrine cells (Kolb 1979), her Fig. 29b), and to assess the probability of coincidental association of puncta (Han and Massey, 2005; Dedek et al., 2006), images of Cx36-fluorescent puncta were flipped horizontally in fields double labeled for Cx45 and Cx36 (Fig. 2B, comparing the same set of images as in Fig. 2A). With the flipped image of Figure 2B2, there was rarely total overlap of individual green and red puncta (Fig. 2B3), compared with nearly total overlap in non- flipped images. Cx45-positive puncta exhibiting an estimated greater than half area of overlap versus less than half area overlap with Cx36-positive puncta represented 5.4% (n = 5; SEM = 0.4%) versus 2.8% (n = 5; SEM = 0.2%), respectively, compared with 87% colocalization in the nonflipped images. This indicates that Cx45/Cx36 colocalization was not simply a consequence of the high density and large size of Cx36 puncta in the IPL.

To confirm that monoclonal and polyclonal anti-Cx36 antibodies generated against different sequences in Cx36 produced the same level of Cx36 detection, rather than potentially differential levels of labeling, thereby affecting the outcome of Cx36/ Cx45 colocalization counts when using these different antibod- ies, sections were double labeled for Cx36 with monoclonal anti- Cx36 combined with polyclonal anti-Cx45. This combination of two anti-Cx36 antibodies showed nearly total overlap of red and green Cx36-immunopositive puncta (Fig. 2C). Similarly, we tested whether the widely used monoclonal anti-Cx45 antibody from Millipore Bioscience Research Reagents and the newly gen- erated polyclonal anti-Cx45 antibody recognized the same or partially distinct subsets of Cx45-puncta in the IPL. Double la- beling with these antibodies also showed nearly total overlap of immunopositive puncta (Fig. 2D).

| Table 2. Quantitative immunofluorescence analysis of Cx45 colocalization relationships with Cx36, ZO-1, ZO-2, and ZONAB in the IPL of adult mouse retina |
|-----------------|-----------------|--------------------|
| Proteins examined | Protein (total puncta counted) | Percentage colocalization |
| Cx45 positive for Cx36 | Cx45 (766) | 46 ± 7.2 |
| Cx45 positive for Cx36 | Cx45 (2952) | 87 ± 1.3 |
| Cx45 positive for ZO-1 | Cx45 (737) | 71 ± 2.8 |
| Cx45 positive for ZO-2 | Cx45 (432) | 7.0 ± 2.1 |
| Cx45 positive for ZONAB | Cx45 (472) | 4.7 ± 0.4 |

Values in parentheses represent the total number of Cx45-positive or Cx36-positive puncta counted in 4–10 separate single-scan confocal images of the IPL in retinas from two to four mice.

Values represent means ± SEM.

Data from mice prepared with 40 ml of fixative, with immediate washout.

Data from mice prepared with 20 ml of fixative, with immediate washout.

FRIL analysis of Cx45 colocalization with Cx36

In samples double labeled for Cx36+Cx45, gap junctions con- taining Cx45 were found in both OFF and ON sublaminae of the IPL. In addition, a few double-labeled gap junctions were found in the outer plexiform layer (data not shown). In 24 FRIL replicas (21 replicas from mice and 3 from rats) that had been double labeled for Cx36+Cx45, a large “N” of 671 immunogold-labeled gap junctions were detected (599 in mouse and 73 in rat; Table 3). Of these, 91% (545 of 599) in mouse IPL and 85% (62 of 73) in rat IPL were labeled for Cx36 only (Fig. 3A2), 8% (48 of 599) and 14% (10 of 73) in mouse and rat, respectively, were double la- beled for both Cx36 and Cx45 (Fig. 3A1, B, C), and 1% (6 of 599) and 1.4% (1 of 73) of those in mouse and rat, respectively, were labeled for Cx45 only (Fig. 3D). Approximately 44% (28 of 64) of Cx45-containing gap junctions were “miniature” gap junctions [i.e., contained <100 connexons; defined by Rash et al. (2007b); see also Rash et al. (1996)], whereas 50% (32 of 64) contained 400 and 2200 connexons (0.2–0.5 μm in diameter; defined as “medium-size”), and 6% (4 of 64) contained between 400 and 2200 connexons (0.2–0.5 μm in diameter; defined as “large”). Overall, 91% (48 of 53) of Cx45-containing gap junctions in mouse IPL were colabeled for Cx36. Double-labeled “mini” gap junctions and those larger gap junctions in which one or the other connexin is present in low amounts (i.e., <30 connexons) may be difficult to detect by FRIL or by conventional immunofluorescence imaging (Kamasawa et al., 2006), regard-
less of the overall size of the gap junction, potentially leading to a slight underestimate of colocalization (tested next).

The ratio of the number of gold beads for Cx45:Cx36 within individual Cx45-containing gap junctions varied from 6:1 (i.e., more Cx45) to 1:36 (i.e., much more Cx36), with overall LEs ranging from LE = 1:3 to LE = 1:59 (i.e., from one gold bead per three connexons to one gold bead per 59 connexons). At the lower LE, some of the few examples of gap junctions containing Cx45 without Cx36 in samples labeled by our standard protocol (1% of total; 6 of 671) may represent “false negatives” for Cx36.

As a test for a threshold of detection for FRIL based on LE = 1:30 versus LE = 1:10, three replicas were labeled for 5 h in primary antibodies and 21.5 h in secondary antibodies. In these replicas, 156 relatively heavily labeled gap junctions were found, 88% (137 of 156) of which were labeled for Cx36 only, 12% (19 of 156) of which were labeled for Cx36 + Cx45 (Fig. 3E), and 0% of which (0 of 156) were labeled for Cx45 only, suggesting that very nearly 100% of Cx45-containing gap junction plaques also contain Cx36, and that no significant population of gap junctions contained only Cx45 without Cx36. The increased ratio of Cx45-containing to Cx36-containing gap junctions after longer labeling times is not attributable to increased nonspecific labeling because almost all of the largest plaque gap junctions (11 of 156) and the only large string gap junction encountered (1 of 156), where nonspecific labeling would be most apparent, were labeled only for Cx36 and not for Cx45. Interestingly, in heavily labeled samples (Fig. 3E), the gold beads were concentrated near the periphery of gap junctions, a distribution that is suggestive of steric hindrance of antibody access to the center of larger gap junctions (investigated further below).

Morphological characteristics of Cx45-containing gap junctions

We previously showed that in IPL, Cx36 was present in a wide variety of morphological classes of gap junctions, designated as crystalline plaque, noncrystalline plaque, reticular, ribbon, and

Figure 3. FRIL localization of Cx45 versus Cx36 in mouse (A–D) and rat (E) IPL. A, Neuronal gap junctions in adjacent cells, one of which (1; 210 connexons) is labeled for both Cx45 (one 18 nm gold and two 6 nm gold beads (arrowheads)) and Cx36 (~450 connexons; eight 12 nm gold beads), and the second of which (2) is labeled for Cx36 only (ten 12 nm gold beads); overall LE = 1:19; 2, LE = 1:45. B, Medium-size plaque gap junction (139 connexons) labeled for Cx45 (three 6 nm gold beads (arrowheads) and one 18 nm gold bead) and Cx36 (five 12 nm gold beads); overall LE = 1:15. C, Medium-size reticular gap junction (106 connexons) labeled for Cx45 (six 6 nm gold beads; arrowheads) and for Cx36 (three 12 nm gold beads). Two gold beads (thin cross-bars) within the apparent “radius of uncertainty” for connexon labelling represent “false-positive” labeling (i.e., “noise”) because stereoscopic analysis (data not shown) revealed that both were >1 μm above the replica on an unstable wisp of undissolved Lexan plastic. Overall LE = 1:12. D, Small gap junction (68 connexons) in rat IPL labeled for Cx45 only (one 18 nm and three 6 nm gold beads; zero 12 nm gold beads are present for Cx36). Cx45 LE = 1:17. This gap junction may be too small to allow positive analysis regarding Cx36. A and C are from same replica. E, Large plaque gap junction (797 connexons) with small area of P-face and larger area of E-face, labeled for both Cx45 (three 12 nm gold beads (arrows) and one 30 nm gold bead) and for Cx36 (seventeen 6 nm gold beads (arrowheads) and forty-one 18 nm gold beads; labeled for approximately twice as long as all other FRIL images shown; see Materials and Methods for details). Overall LE = 1:13. Scale bars, 0.1 μm.
Figure 4. Eight generalized connexin/connexon configurations proposed to occur within neuronal gap junctions. A, B, Homomeric heterotypic Cx36-to-Cx36 channels (A) and Cx45-to-Cx45 channels (B). C. Cx36 in the top cell and Cx45 in the bottom cell, with Cx36 linking to Cx45 (“homomeric heterotypic”). D. Cx43 in the top cell and Cx45 in the bottom cell, with Cx43 linking to Cx45 (“homomeric heterotypic”). E, Both Cx36 and Cx45 in both cells, with Cx36 linked only to Cx36 and Cx45 linked only to Cx45 (“homomeric bimorphy”). F. Both Cx36 and Cx45 in both cells, with both connexons on one side linked to both connexons on the other side (“homomeric bimorphy”). G, Cx36 only in the top cells linking to connexons containing a mixture of Cx36/Cx45 in the bottom cell (homomeric to heteromeric channels). H, Each connexon contains both Cx36 and Cx45 (heteromeric connexons), linking to heteromorphy channels in the other cell (heteromeric to heteromeric channels).

string configurations, with string gap junctions associated with more abundant ZO-2 labeling and plaque gap junctions associated with more abundant ZO-1 labeling (Ciolofan et al., 2006; Kamasawa et al., 2006). Ribbon and string configurations represented ~50% of Cx36-containing gap junctions in the OFF sublamina. In contrast, almost all (94%; 60 of 64) Cx45-containing gap junctions in this study were plaques (Fig. 3A–C), and several (6%; 4 of 64) were reticular gap junctions (i.e., plaques with internal voids) (Fig. 3D), but none (0%; 0 of 64) were string or ribbon gap junctions. These differences in morphology suggest that Cx45-containing gap junctions may be associated with different scaffolding proteins than those containing only Cx36 (see below).

Absence of heterotypic Cx36/Cx45 gap junctions in HeLa cells

Teubner et al. (2000) reported that in HeLa cells separately expressing Cx36 and Cx45, “None of these co-cultures demonstrated transfer above background of either Neurobiotin or Lucifer yellow (data not shown)”, and concluded that Cx45 does not form functional couplings with Cx36. To investigate conditions under which Cx45 and Cx36 engage in gap junction plaque formation together and separately, we used HeLa cells expressing Cx45 and Cx36 to evaluate several potential connexin coupling combinations by immunofluorescence imaging, which was not done in the previous studies. In principal, a connexon hemichannel may assemble from six identical connexin monomers to form “homomeric” connexons (Fig. 4A–C). In turn, connexons may link to identical homomeric connexons to form “homotypic” channels (Fig. 4A,B,E), or they may link to connexons containing unlike connexins to form “heterotypic” channels (Fig. 4C,D,F). Because FRIL revealed that Cx45 was almost always accompanied by Cx36 within individual hemi-plaques in retina (supporting one of Fig. 4E–H), we tested whether homomeric Cx45 hemichannels can form heterotypic plaques by coupling with homomeric Cx36 hemichannels (as in Fig. 4C). First, as a positive control for the ability of HeLa cells to form heterotypic channels, we used FRIL to show that Cx45 expressed in HeLa cells formed large numbers of Cx45-foci (Fig. 4A), indicating the ability of HeLa cells to traffic coexpressed connexins to the same membrane locations, apparently to the same gap junctional plaques, as occurs in retina. Also encountered were pairs of adjacent Cx36-HeLa cells that were each transiently transfected with Cx45 and that formed Cx36/Cx45 colocalized plaques at points of plasma membrane apposition (Fig. 5C), indicating the ability of HeLa cells to traffic the coexpressed connexins to the same membrane locations, apparently to the same gap junctional plaques, as occurs in retina. Also encountered were pairs of adjacent Cx36-HeLa cells that were each transiently transfected with Cx45 and that formed Cx36/Cx45 colocalized plaques at points of plasma membrane apposition (Fig. 5C), indicating the ability of HeLa cells to traffic the coexpressed connexins to the same membrane locations, apparently to the same gap junctional plaques, as occurs in retina. Also encountered were pairs of adjacent Cx36-HeLa cells that were each transiently transfected with Cx45 and that formed Cx36/Cx45 colocalized plaques at points of plasma membrane apposition (Fig. 5C), indicating the ability of HeLa cells to traffic the coexpressed connexins to the same membrane locations, apparently to the same gap junctional plaques, as occurs in retina. Also encountered were pairs of adjacent Cx36-HeLa cells that were each transiently transfected with Cx45 and that formed Cx36/Cx45 colocalized plaques at points of plasma membrane apposition (Fig. 5C), indicating the ability of HeLa cells to traffic the coexpressed connexins to the same membrane locations, apparently to the same gap junctional plaques, as occurs in retina. Also encountered were pairs of adjacent Cx36-HeLa cells that were each transiently transfected with Cx45 and that formed Cx36/Cx45 colocalized plaques at points of plasma membrane apposition (Fig. 5C), indicating the ability of HeLa cells to traffic the coexpressed connexins to the same membrane locations, apparently to the same gap junctional plaques, as occurs in retina. Also encountered were pairs of adjacent Cx36-HeLa cells that were each transiently transfected with Cx45 and that formed Cx36/Cx45 colocalized plaques at points of plasma membrane apposition (Fig. 5C), indicating the ability of HeLa cells to traffic the coexpressed connexins to the same membrane locations, apparently to the same gap junctional plaques, as occurs in retina. Also encountered were pairs of adjacent Cx36-HeLa cells that were each transiently transfected with Cx45 and that formed Cx36/Cx45 colocalized plaques at points of plasma membrane apposition (Fig. 5C), indicating the ability of HeLa cells to traffic the coexpressed connexins to the same membrane locations, apparently to the same gap junctional plaques, as occurs in retina. Also encountered were pairs of adjacent Cx36-HeLa cells that were each transiently transfected with Cx45 and that formed Cx36/Cx45 colocalized plaques at points of plasma membrane apposition (Fig. 5C), indicating the ability of HeLa cells to traffic the coexpressed connexins to the same membrane locations, apparently to the same gap junctional plaques, as occurs in retina. Also encountered were pairs of adjacent Cx36-HeLa cells that were each transiently transfected with Cx45 and that formed Cx36/Cx45 colocalized plaques at points of plasma membrane apposition (Fig. 5C), indicating the ability of HeLa cells to traffic the coexpressed connexins to the same membrane locations, apparently to the same gap junctional plaques, as occurs in retina.
incapable of forming heteromeric connexins (Evans and Martin, 2002). Nevertheless, the arrangement of homomeric versus heteromeric connexons might be ascertained by double-replica FRIL if labeling for the two connexins were to be unequivocally segregated (examined in next section).

Double-replica analysis of Cx45 and Cx36 in apposed hemiplaques in rat IPL
Because conventional FRIL is based on random fracturing through complex tissue, a sufficiently large sample size will reveal all possible configurations of gap junctions and accurately reveal the range of connexon labeling and connexin pairing relationships. The absence of a significant population of gap junction plaques containing only Cx45 (0.6%, all data combined) means that there is no significant population of gap junctions containing only Cx45 in only one hemiplaque or in both apposed hemiplaques. Only by the use of double replicas showing both sides of the same gap junction could it be proven that both connexins are contained in both sides of a pair of hemiplaques. Moreover, the above immunofluorescence results documenting gap junction plaque-forming capability of Cx45 and Cx36 in HeLa cell expression systems underscored the necessity for examining matched double replicas of Cx45-containing gap junctions in retina.

In both single and double replicas, the fracture plane always separates apposed connexons at their point of contact in the extracellular space (Fig. 6A) [as previously shown (Fujimoto, 1997; Rash and Yasumura, 1999; Nagy et al., 2004)], separating gap junctions into two complementary hemiplaques, each containing connexons from only one or the other cell (Fig. 6B, C). Immunogold labeling of cytoplasmic epitopes of connexons occurs in the residual cytoplasm beneath the Pt/C replica (Fig. 6B, C). Moreover, labeling is approximately equal beneath both E-face pits and P-face particles, demonstrating that the connexons of the subjacent cell are retained and not washed away during SDS cleaning (Fujimoto, 1997; Rash and Yasumura, 1999). In the case of double replicas, however, both fracture faces are retained and replicated, and both sides are approximately equally well labeled in the matching complementary images of both E-face pits and P-face IMPs (Fig. 6B–G). This feature for clean separation of hemiplaques in vertebrate gap junctions allows definitive assignment of one or multiple connexons to individual hemiplaques belonging to one or the other apposed cell (Rash et al., 2001, 2004; Nagy et al., 2004; Kamasawa et al., 2005), and equally important, potentially allows assessment of whether Cx45 and Cx36 are intermixed (as heteromeric connexons) (Fig. 4G, H) or segregated into domains of homomeric connexons (Fig. 4E).

In double replicas of complex neuropil, multiple structural markers are used to find the same gap junction in both complementary replicas. However, corresponding areas in one or the other replica are often missing because (1) one or the other may be covered by grid bars, even when using thin-bar slot grids (Fig. 6D, D'; dotted lines on each sample correspond to locations of obscuring grid bars in the complementary replica); (2) fragmentation and loss of complementary areas may occur during sample processing (Fig. 6D, D'; outlines indicate portions lost in its complementary replica); or (3) one or the other matching area was obscured by adhering debris (one example, not shown). Nevertheless, in a large portion of one set of matched double replicas

Figure 5. Double-immunofluorescence labeling of connexins in cultures containing mixtures of HeLa cells expressing Cx45, Cx43, or Cx36. A, Coculture of cells stably expressing Cx45 with those expressing Cx43, showing colocalization of Cx45/Cx43 at abutments between Cx45-HeLa and Cx43-HeLa cells (A1–A3, arrows, yellow puncta in A3). B, Coculture of cells expressing Cx45 with those expressing Cx36, showing Cx45-positive puncta at cell appositions among colonies of Cx45-HeLa cells (B1, bottom right half of field and arrows) and Cx36-positive puncta at appositions among Cx36-HeLa cells (B2, top left half of field and arrowhead). There is a lack of Cx45/Cx36 colocalization at abutments between Cx45-HeLa and Cx36 HeLa cells (B3, double-headed arrows). C, HeLa cells stably expressing Cx36 and transiently transfected with Cx45, showing two apposed cells (arrows) expressing both connexins, and Cx45/Cx36 colocalization at points of cell–cell contact (arrow). D, Two HeLa cells stably expressing Cx36 and transiently transfected with Cx45, showing Cx45 localized intracellularly (D1, double-headed arrows), Cx36/Cx45 colocalization at points of cell–cell contact between cells expressing both connexins (arrowhead), and absence of Cx45 at points at which double-expressing cells make contacts with cells expressing only Cx36 (arrow). Scale bars, 10 μm.
Figure 6. Images and the interpretation of labeling in matched double replicas. A–C, Scale diagrams showing gap junction before and after freeze fracturing and retrieval of complementary replicas. A, The fracture plane (dashed line) separates connexons at their points of contact in the extracellular space, leaving all connexons of the bottom cell for subsequent immunogold labeling. B, C, Creation of double replicas having complementarity of E- and P-fracture faces. Each matching replica contains connexons from only one cell, but the E-face of one cell and the P-face of the other cell. D–D′, Images from dissecting microscope showing matched double replicas containing three strips of retina after SDS washing and immunogold labeling. E, Low-magnification image of one side of a complementary replica, with locations of two of the mapped gap junctions indicated by arrows. Ax, Axons. F, F′, G, G′, High-magnification stereoscopic views of complementary replicas of two gap junctions, with 10 nm gold labeling Cx36 and 5 nm gold labeling Cx45. Cx36 labeling is present beneath three quadrants of both gap junctions (E-face pits in top replica and complementary P-face particles in bottom replica). In the bottom right quadrant, 5 nm Cx45 labels are present. Initial precoating with 2–3 nm of carbon partially obscured E-face connexon pits. The arrow (in F) indicates a gold bead beyond the 28 nm radius of immunogold labeling. The arrowhead (in G′) indicates a 5 nm gold label for Cx45, with possible corresponding label in the complementary image (G, arrow). Scale bars, 0.1 μm unless otherwise indicated.
of rod and cone bipolar cells. The outer oval delineates the area with increased Cx45 labeling (twelve and eleven 5 nm gold beads, \( A \ vs \ B \)), and the inner oval delineates the domain of Cx36 labeling (eleven and nine 10 nm gold beads, \( A \ vs \ B \)). The inner oval delineates the domain of Cx36 labeling (eleven and nine 10 nm gold beads, \( A \ vs \ B \)).

Figure 7. Cx45/Cx36 double-labeled gap junction adjacent to a ribbon of E-face particles/P-face pits (rectangles) that are found postsynaptic to ribbon synapses, which in IPL, are characteristic of rod and cone bipolar cells. The outer oval delineates the area with increased Cx45 labeling (twelve and eleven 5 nm gold beads, \( A \ vs \ B \)), and the inner oval delineates the domain of Cx36 labeling (eleven and nine 10 nm gold beads, \( A \ vs \ B \)).

(Fig. 6D, D’), we found >160 immunogold-labeled gap junctions, including 11 Cx45-containing gap junctions that had matching complements, five of which are illustrated here (Figs. 6, 7; supplemental Fig. S1, available at www.jneurosci.org as supplemental material). Eight of these 11 pairs were labeled for both Cx45 and Cx36 in both matching hemiplaques. In these matched pairs, the number of gold beads for the two connexins was substantially greater in one versus the other hemiplaque in six pairs (Fig. 6F, G; supplemental Fig. S1A, available at www.jneurosci.org as supplemental material), and approximately equal in two paired hemiplaques (Fig. 7). In six of eight pairs, Cx45 and Cx36 appeared to be segregated into domains, with labels for both connexins beneath their respective complementary sectors in the mirror hemiplaques (i.e., Cx45 opposite Cx45 and Cx36 opposite Cx36) (Figs. 6F, G, 7). This observed segregation of labels implies that the connexons are present in homomorphic connexons that are further segregated into domains containing either Cx36 or Cx45 but not both (i.e., correspond to Fig. 4E but not to Fig. 4G or 4H). Variations in number of labels for each connexin in matched pairs are attributed to stochastic differences arising from the numerically low labeling efficiency (LE = 1:15 to 1:20) in these matched pairs versus the relatively few connexons of each connexin type, particularly in smaller gap junctions. Minor discrepancies in positions of gold beads (Fig. 6G, arrows) are attributed to the 28 nm “radius of uncertainty” of immunogold labeling (Fig. 6B, C) or to the displacement of connexons into lipid “blebs” formed during SDS washing (Rash and Yasumura, 1999; Kamasawa et al., 2006). These few examples of apparent segregation of each connexin into discrete domains do not eliminate the possibility that homomeric homotypic Cx45- and Cx36-connexons may also be intermixed in some gap junctions. However, the relatively low labeling efficiency could not preclude the existence of a few heteromeric connexons. Nevertheless, our immunocytochemical data showing absence of Cx45/Cx36 homotypic coupling (Fig. 5), plus FRIL data in which 60 of 64 (and 19 of 19; Table 3) Cx45-labeled gap junctions also contained Cx36, combined with data from six of eight mirrored gap junctions showing approximate alignment of Cx45- versus Cx36-containing domains, unequivocally document that Cx45 and Cx36 are present, together on both sides of the same gap junction, presumably with homomorphic Cx45 connexons coupling with homomorphic Cx36 connexons, and homomorphic Cx45 connexons coupling with homomorphic Cx36 connexons. This bihomotypic coupling pattern occurred in most if not all Cx45-containing gap junctions in mouse and rat IPL.

In three of the eight mirror pairs from the ON sublamina of IPL, double-labeled gap junctions were adjacent to linear arrays of postsynaptic E-face particles characteristic of ribbon synapses (Fig. 7, inscribed box), which in IPL, occur only in cells postsynaptic to rod and cone bipolar cells (Kolb, 1979). In each of these three mirror pairs, the cell that was postsynaptic to the ribbon synapse was linked to an unidentified third cell by a gap junction containing both Cx45 and Cx36 in both sides (inscribed oval). Thus, these coupled cell pairs that were postsynaptic to bipolar cells in the ON sublamina coexpressed and cotargeted Cx45 and Cx36.

Of the remaining three gap junction pairs with unequal labeling of apposed hemiplaques (from 11 total), one small gap junction (37 connexons) was labeled for both Cx36 (seven gold beads) and Cx45 (two gold beads) on one side but was not labeled for either connexin on its matched complement (not shown). Assuming that the gap junction contained no more than ~10 Cx45 connexons and 27 Cx36 connexons, and a nominal LE = 1:20, the complete absence of labeling for either connexin on the unlabeled side would be consistent with normal stochastic variability of labeling. The second asymmetrically labeled gap junction (~100 connexons) had three gold beads for Cx45 and seven gold beads for Cx36 on one side and zero gold beads for Cx45 and five gold beads for Cx36 on the complementary surface. The third of these unequally labeled pairs was a large gap junction (>400 connexons) that was well labeled for Cx36 in both the upper and lower hemiplaques (37 and 32 gold beads, respectively) (supplemental Fig. S1B, available at www.jneurosci.org as supplemental material) but contained only a single gold bead for Cx45 on one side.
and none on the other side. The latter pair of images could imply that there were equally few Cx45 connexons in the mirror hemiplaque, or it could imply the presence of a few unpaired Cx45 connexons. However, this discrepancy of one versus zero gold beads for Cx45 is also consistent with normal stochastic differences in connexin labeling efficiency.

### Statistical analysis of Cx45 relationships with Cx36 in retinal neurons and gap junctions

To test various hypotheses regarding expression and localization of Cx36 and Cx45 in the retina, we used a web-based on-line statistical analysis system to calculate confidence intervals (http://www.measuringusability.com/wald.htm) (as described in Hypothesis 3).

**Hypothesis 1: Cx45 and Cx36 are separately expressed in a neuron subtype-specific manner**

The hypothesis that Cx45 is not coexpressed in cells that express Cx36 is nullified by finding in single replicas that most (60 of 64) Cx45-containing gap junctions also had Cx36 labeling (Table 3), and further nullified by 19 of 22 mirror hemiplaques in double replicas showing that both connexins occur in the same cell (Figs. 6, 7). However, the four of 64 examples of Cx45 labeling without Cx36 labeling are consistent with one of the following: (1) incomplete (i.e., false-negative) labeling for Cx36 in a very small fraction (4 of 64) of Cx45 expressing cells, (2) lack of simultaneous insertion of Cx36 with Cx45 into plaques [whereas endocytotic retrieval of connexons involves internalization of structurally coupled membrane patches (Gaietta et al., 2002)], or (3) existence of a small fraction of Cx45-expressing cells that do not coexpress Cx36. In the latter case, Hypothesis 1 might still apply, but at most, only to a very small fraction of Cx45-expressing cells.

**Hypothesis 2: Cx45-expressing cells have hemiplaques containing only Cx45, either as monohomotypic (Cx45-to-Cx45) or heterotypic (Cx45-to-Cx36) gap junctions**

Hypothesis 2 is invalidated by the demonstration in single replicas that >90% (60 of 64) of Cx45-containing hemiplaques also contain Cx36 (Fig. 3, Table 3). Similarly, double replicas showed that none of 11 matched pairs of Cx45-containing gap junctions had only Cx45 in either of their apposed hemiplaques (Figs. 6, 7; supplemental Fig. S1, available at www.jneurosci.org as supplemental material), further invalidating Hypothesis 2.

**Hypothesis 3: Cx36 is present in Cx45-containing gap junctions, forming bimolecular plaques**

Table 4 presents the percentages, along with 95% confidence intervals, for Cx45+/Cx36 forming bimolecular gap junctions in mouse and rat IPL in both single replicas and double replicas, where 93.8 to 100% of Cx45-containing gap junctions also contain Cx36. Moreover, eight of 11 pairs of Cx45-containing gap junctions found in double replicas contained both Cx45 and Cx36 in both sides, with six of those pairs, to the limit of the radius of uncertainty of immunogold labeling, having near-precise mirror distributions of labels for both connexins, thereby demonstrating the presence of bimolecular gap junctions that also are primarily or exclusively homomeric.

### Cx45 and ZO-1 in retina of Cx36 ko mice

The presence of ZO-1 at gap junctions containing Cx45 suggested its association either with Cx45 or exclusively with Cx36, which is also contained in most of these junctions and with which ZO-1 was recently found to be highly colocalized (Li et al., 2004a). This question was addressed by comparing labeling patterns of Cx45 in retina of wild-type and Cx36 ko mice. Double-immunofluorescence labeling was conducted to determine whether the presence of Cx36 at Cx45/ Cx45 colocalized puncta was required for Cx45/ZO-1 association.
As shown by image overlay (Fig. 8E), substantial Cx45/ZO-1 overlap was found in Cx36 ko mice. Counts of puncta revealed that 78 ± 4.2% and 78 ± 2.4% of Cx45-puncta were colocalized with ZO-1 in the IPL of retina from wild-type and Cx36 ko mice, respectively, which indicated that elimination of Cx36 had little affect on Cx45/ZO-1 association and suggested direct molecular interaction of Cx45 with ZO-1, as examined further, below. Confocal double immunofluorescence in vertical sections of retina from wild-type C57BL/6 mice indicated similar labeling and distribution of Cx36 and Cx45 in the IPL (Fig. 8F,G) as observed in CD1 mice (Figs. 1, 2). Double-immunofluorescence imaging in retina of Cx36 ko mice showed an absence of labeling for Cx36 in the IPL (Fig. 8H) and reduced labeling of Cx45 (Fig. 8I). Counts of Cx45-puncta in eight randomly selected fields in the IPL from wild-type and Cx36 ko mice revealed a significant 25% reduction (p < 0.05) of Cx45-puncta in the ko mice.

FRIL analysis of Cx45 colocalization with ZO-1
SDS digestion (as used in SDS-FRL and FRIL) does not separate transmembrane proteins from many of their tightly bound cytoplasmic scaffolding and accessory proteins, leaving many of those molecules available for detection by immunogold labeling (Fujimoto, 1997; Fujimoto et al., 1997; Ciolofan et al., 2006). In a replica that was double labeled for Cx45 (Millipore Bioscience Research Reagents monoclonal antibody) plus ZO-1 (Invitrogen rabbit polyclonal antibody), 48 labeled gap junctions were found (Fig. 9). Of these, 44% (21 of 48) were double labeled for Cx45 plus ZO-1 (Fig. 9A,B), 4% (2 of 48) were labeled for Cx45 only (data not shown), and 52% (25 of 48) were labeled for ZO-1 only (Fig. 9C,D), including five string gap junctions (Fig. 9C). In this replica, which was labeled relatively briefly (1.25 h in primary antibodies and 16 h in secondary antibodies), 91% (21 of 23) of Cx45-labeled gap junctions were colabeled for ZO-1. This means that ZO-1 is on both sides of most, if not all, Cx45-containing gap junctions. The remaining 25 gap junctions (52%) were labeled for ZO-1 without Cx45, consistent with observations from immunofluorescence labeling (see above) and from FRIL (Ciolofan et al., 2006; Kamasawa et al., 2006) that many from the larger pool of homotypic Cx36-containing gap junctions also have closely bound ZO-1. With respect to ultrastructural correlates, almost all of those ZO-1/Cx45 double-labeled gap junctions were of the plaque type, and none were strings or ribbons. In contrast, ~50% of Cx36-containing gap junctions in the OFF sublamina were string and ribbon configurations, and those were shown to be relatively deficient in ZO-1 (Ciolofan et al., 2006; Kamasawa et al., 2006).
Interaction of Cx45 with ZO-1 in HeLa cells and retina

Cultured HeLa cells stably transfected with Cx45 were used to document Cx45 association with ZO-1. In control empty-vector-transfected cells, labeling of ZO-1 typically appeared as fine, intermittent puncta or as short, continuous strands of immunofluorescence at cell–cell contacts (Fig. 10A1), whereas Cx45 immunoreactivity was entirely absent (Fig. 10A2, A3). In Cx45-HeLa cells, labeling of ZO-1 had a more dispersed appearance and was localized both at cell–cell contacts and intracellularly (Fig. 10B1). Nearly all cells displayed labeling of Cx45 in puncta...
at contacts between cells and/or intracellularly (Fig. 10B2). Double-immunofluorescence labeling indicated partial Cx45/ZO-1 overlap at both of these subcellular locations (Fig. 10B3). Laser-scanning confocal double-immunofluorescence imaging further revealed that individual ZO-1-positive puncta at cell margins were immunopositive for Cx45 (Fig. 10C).

Immunoblotting and coimmunoprecipitation were used to confirm anti-Cx45 antibody specificity and to examine molecular interaction of Cx45 with ZO-1. In lysates of Cx45-HeLa cells examined using 12.5% gels, monoclonal anti-Cx45 (Millipore Bioscience Research Reagents) detected a band migrating at ~45 kDa (Fig. 11A, lane 2), whereas this band was absent in empty vector-transfected HeLa cells (Fig. 11A, lane 1). A similar comigrating band was detected in homogenates of retina (Fig. 11A, lane 4), but was absent in homogenates of liver (Fig. 11A, lane 3), which was used as a negative control. After IP of ZO-1 from homogenates of retina using polyclonal anti-ZO-1, IP material probed with monoclonal anti-Cx45 was found to contain Cx45 (Fig. 11A, lane 6) comigrating with corresponding bands detected with monoclonal anti-Cx45 in retina (5% of input protein in IP) and in Cx45-HeLa cells. Cx45 was not detectable in immunoblotted material after omission of anti-ZO-1 in the IP procedure (Fig. 11A, lane 5). A separate blot again shows a Cx45 band in retina (Fig. 11A, lane 7) and absence of this band after IP of ZO-1 from retina using anti-FLAG IgG as control antibody (Fig. 11A, lane 8). By Western blotting, we have previously shown the presence of ZO-1 in retinal homogenates used as input for IP of ZO-1 (Li et al., 2004a). Immunoblotting of Cx45-HeLa cells using 12.5% gels and probing of membranes with polyclonal anti-Cx45 gave a similar detection of Cx45 (Fig. 11B, lane 1), which was absent in empty vector-transfected cells (Fig. 11B, lane 2). In 10% SDS gels, the polyclonal anti-Cx45 detected a doublet of bands (Fig. 11C), corresponding in molecular weight to previously reported phosphorylated and nonphosphorylated forms of Cx45 (van Veen et al., 2000). The polyclonal anti-Cx45 also detected Cx45 obtained from retina after IP of Cx45 with monoclo-
These data demonstrated similar labeling specificity for all Cx45 antibodies tested.

Homogenates of mouse retina were used with fusion proteins containing GST linked separately to each of the three PDZ domains in ZO-1 for in vitro pull-down assays to examine direct binding of Cx45 to the PDZ domains of ZO-1. As shown by immunoblotting (Fig. 12A), eluates of protein from beads coupled to either the GST-PDZ1 (Fig. 12A, lane 2) or GST-PDZ3 (Fig. 12A, lane 4) domain of ZO-1 were devoid of Cx45, whereas protein eluted from beads coupled to the GST-PDZ2 domain of ZO-1 (Fig. 12A, lane 3) contained Cx45, which migrated as a band corresponding to Cx45 detected in homogenates of mouse retina (Fig. 12A, lane 1). Stripping of the blot in Figure 12A and reprobing with anti-GST antibody revealed equal loading of the three PDZ domain fusion proteins (Fig. 12B). As a positive control for the functional binding capacity of the PDZ1 domain fusion protein, we have previously shown that this fusion protein

![Figure 11. Immunoblots demonstrating co-IP of Cx45/ZO-1 from retina and detection of Cx45 with polyclonal anti-Cx45. A, Cx45 detection by monoclonal anti-Cx45 in lysates of Cx45-transfected HeLa cells (lane 2) and in homogenate of mouse retina (lanes 4 and 7), but not in empty vector-transfected HeLa cells (lane 1) or in liver (lane 3). After IP of ZO-1 from retina homogenate using polyclonal anti-ZO-1, monoclonal anti-Cx45 detects Cx45 (lane 6) comigrating with Cx45 from retina homogenate (lane 4) and from Cx45-transfected HeLa cells (lane 2), but not in IP material after omission of anti-ZO-1 during the IP procedure (lane 5) or after IP of ZO-1 from retina using anti-FLAG as control IgG (lane 8). B, C, Polyclonal anti-Cx45 detection of Cx45 as a single band using 12.5% SDS-gel (lane 1) and as a doublet of bands using 10% SDS-gel (C, lane 1) in lysates of Cx45-transfected HeLa cells, and absence of these bands in empty vector-transfected HeLa cells (B, C, lane 2). D, IP of Cx45 from retina with monoclonal anti-Cx45, showing detection of a Cx45 band with polyclonal anti-Cx45 in IP material (lane 3), corresponding to a Cx45 band detected by the polyclonal antibody in lysate from Cx45-transfected HeLa cells (lane 2), and absence of the band in empty vector-transfected HeLa controls (lane 1) and after omission of monoclonal anti-Cx45 during the IP procedure (lane 4).]

![Figure 12. Immunoblots demonstrating Cx45 pull-down with the PDZ2 domain of ZO-1. Lysates of Cx45-transfected HeLa cells were incubated with GST fusion proteins containing the PDZ1, PDZ2, or PDZ3 domain of ZO-1. A, Immunoblots of pull-down proteins probed with monoclonal anti-Cx45 show detection of Cx45 after pull-down with the PDZ2 domain of ZO-1 (lane 3), but not after pull-down with the PDZ1 or PDZ3 domain of ZO-1 (lane 2 and 4). The Cx45 band in lane 3 corresponds to detection of the same band in homogenate of retina (lane 1). B, Blot in A was stripped and reprobed with anti-GST antibody to show presence and equal loading of PDZ domain fusion proteins.](image-url)
Co-IP of Cx36 with Cx45 from retina and HeLa cells

Direct interaction of Cx36 with the PDZ1 domain of ZO-1 and interaction of Cx45 with the PDZ2 domain of ZO-1, together with the presence of these two connexins in the same gap junctional hemiplaques, raised the possibility that these two connexins may be linked together by ZO-1 in a tripartite complex. Molecular association of Cx36 with Cx45 was therefore investigated by co-IP of these connexins from mouse retina and from HeLa cells cotransfected with either full-length Cx36 and Cx45 or with full-length Cx45 and truncated Cx36 (Cx36-tr) lacking the C terminus PDZ1 domain-binding motif. As shown in immunoblots probed with polyclonal anti-Cx36, IP of Cx45 with monoclonal anti-Cx45 resulted in detection of Cx36 in retinal IP material (Fig. 13A, lane 2) and in IP material from HeLa cells cotransfected with Cx36/Cx45 (Fig. 13A, lane 1). In contrast, Cx36 was not detected in IP material from HeLa cells cotransfected with Cx45 and truncated Cx36 (Fig. 13A, lane 3), nor was it detected in IP material after omission of anti-Cx45 during the IP procedure (Fig. 12A, lane 4). We have previously shown by Western blotting that truncation of Cx36 by four C-terminal amino acids still allows Cx36 recognition by polyclonal anti-Cx36 antibody (Li et al., 2004a). Similar results were obtained using the reverse IP approach; IP of Cx45 with polyclonal anti-Cx45 resulted in detection of Cx45 in retinal IP material when probed with monoclonal anti-Cx45 (Fig. 13B, lane 2) and in IP material from HeLa cells cotransfected with Cx36/Cx45 (Fig. 13B, lane 1). However, Cx45 was not detected in IP material from HeLa cells cotransfected with Cx45 and truncated Cx36 (Fig. 13B, lane 3) or in IP material after omission of anti-Cx45 during the IP procedure (Fig. 12B, lane 4). Additional control procedures are shown in supplemental Figure S2 (available at www.jneurosci.org as supplemental material). Specifically, input levels of Cx36 and Cx45 are demonstrated in lysates (5%) of Cx36/Cx45-transfected and Cx36-tr/Cx45-transfected HeLa cells and retinal homogenates used for co-IP, and absence of Cx45 or Cx36 in IP material is shown after IP from each of the tissues using monoclonal and polyclonal anti-FLAG antibodies as control IgG.

Discussion

Establishing neuronal gap junctional coupling relationships in the retina is essential for understanding how rod scotopic and mesopic information is “piggybacked” onto the photopic cone pathway (Kolb, 1979; Strettoi et al., 1992; Vaney, 2002). Key to this goal is identification of connexin expression profiles of IPL neurons and determination of the connexin constituents of their gap junctions. A major new finding from FRIL and from SDS-FRL double replicas, and one entirely unexpected based on current literature descriptions of neuron subtype-specific expression of Cx36 versus Cx45 in the IPL of rodent retina, is that Cx36 was nearly always colocalized with Cx45 in Cx45-containing freeze-fractured neuronal gap junctions. If a significant fraction of retinal gap junctions in IPL were to express only Cx45 in homotypic or in heterotypic hemiplaques, as proposed by others, FRIL would have revealed primarily or exclusively Cx45-single-labeled hemiplaques. This was not observed. Suggestions that Cx45 does not form functional heterotypic couplings with Cx36 (Teubner et al., 2000) were further supported by our coexpression experiments, which showed absence of heterotypic puncta in HeLa cells expressing Cx45 and Cx36. Overall, our finding that 87% of Cx45-immunopositive puncta in the IPL were also immunopositive for Cx36 supports the extent of Cx36/Cx45 colocalization across the IPL. Finally, Cx45 and Cx36 often had segregated but mirror distributions in both sides of opposing hemiplaques, forming definitive bihomotypic rather than heterotypic gap junctions, and moreover, that Cx45 and Cx36 were primarily or exclusively in homomeric connexons. These results raise several issues regarding current literature on connexin biology in the IPL.

First, Cx45-positive immunofluorescent puncta in the IPL were reported never to overlap (0%) with Cx36-positive puncta (Han and Massey, 2005) or to exhibit 30% colocalization (Dedek et al., 2006), both contrasting with 87% immunofluorescence colocalization found here. Likewise, the OFF sublamina was reported to be devoid of Cx45 immunofluorescence (Güldenagel et al., 2004a, b).
al., 2000) or to contain a very low density of Cx45 (Maxeiner et al., 2005), whereas we observed relatively abundant Cx45 puncta in the OFF sublamina. Also contradictory, Feingenspan et al. (2001) noted that the heterologous gap junctions between AII amacrine cells and ON cone bipolar cells “appear to be heterotypic because ON cone bipolar cells do not express Cx36,” whereas Deans et al. (2002) stated that “Cx36 is expressed by ON CBs coupled to AII amacrine cells.” In contrast, Maxeiner et al. (2005) stated that “all four classes of OFF cone bipolar cells express Cx45,” and acknowledged “the possibility that both Cx36 and Cx45 are expressed by the same bipolar cell.” We attribute these differences in immunofluorescence detection to fixation-dependent reduction in antibody labeling, as evidenced by a lower percentage of Cx36/Cx45 colocalization after increased fixation (45 vs 87%).

Second, reporter-based approaches to cell-type identification of connexin expression in IPL suggested that neurons expressing Cx45 represent entirely different populations than those expressing Cx36. Thus, Dedek et al. (2006) attributed their 30% colocalization of Cx36 with Cx45 to each connexin exclusively in opposite sides of opposed hemiplaques, which they proposed to represent heterotypic Cx45-cone bipolar to Cx36-AII amacrine cell gap junctions. Although Cx45 lacks dye-coupling competence with Cx36 in HeLa cells transfected with these connexins (Teubner et al., 2000), it was suggested that HeLa cells might lack essential accessory proteins that otherwise permit the hypothesized functional Cx36/Cx45 coupling in IPL neurons (Maxeiner et al., 2005). Because Teubner et al. (2000) had not assayed for connexin colocalization, we investigated similar connexin-transfected HeLa cell lines and showed that they form immunopositive plaques containing homotypic Cx36/Cx36, Cx43/Cx43, and Cx45/Cx45 puncta, as well as heterotypic Cx43/Cx45 puncta at cell–cell abutments, but did not form heterotypic Cx45/Cx45 puncta. Instead, immunofluorescence colocalization of Cx45+/Cx36 in HeLa cell contacts required that both cells simultaneously coexpress both proteins. Because HeLa cells contain requisite scaffolding and targeting proteins to allow separate formation of Cx45-containing and Cx36-containing gap junctions, as well as Cx45+/Cx36 bimhomotypic gap junctions, Cx45 appears unable to form heterotypic puncta with Cx36 in these cells.

Third, our combined results imply that Cx45 and Cx36 are transported to and stabilized in retinal gap junctions by common targeting and scaffolding proteins. Cx36 binds to the PDZ1 domain of ZO-1 (Li et al., 2004a,b), and here we showed Cx45/ZO-1 coassociation in mouse retina and interaction of Cx45 with the PDZ2 domain, which had not been determined previously (Kausalya et al., 2001; Laing et al., 2001; Thomas et al., 2002). Significantly, Cx45/ZO-1 colocalization persisted in IPL from Cx36 ko mice, indicating that this colocalization was not simply attributable to ZO-1 association with Cx36 at Cx45-containing gap junctions. Presence of all three proteins in IPL gap junctions and their co-IP suggest an essential targeting and scaffolding role for ZO-1 in Cx45/Cx36-containing gap junctions. With nine members of the connexin family of proteins reported to associate with ZO-1 (Giepmans, 2004; Li et al., 2004a,b,c; Penes et al., 2005), a regulatory role for ZO-1 at gap junctions is emerging (Li et al., 2004a,b,c; Hunter et al., 2005; Laing et al., 2005; Akoyev and Takemoto, 2007). Binding of Cx36 and Cx45 to different ZO-1 PDZ domains suggests that ZO-1 may simultaneously interact with Cx36 and Cx45 in a tripartite manner, thereby tethering the two connexins within gap junctions. A critical role of ZO-1 in mediating this association was indicated by abolition of Cx36/Cx45 co-IP after removal of the C-terminal 4 aa residues of Cx43 (this report), which are required for the interaction of Cx36 with the PDZ1 domain of ZO-1 (Li et al., 2004a).

Fourth, heterotypic gap junctional coupling between Cx36-expressing AII amacrine cells and Cx45-expressing ON-cone bipolar cells was advanced as a key element in the rod pathway, based on substantial alteration of electroretinograms in Cx45 ko mice (Maxeiner et al., 2005). We found that absence of Cx36 in Cx36 ko mice was accompanied by 25% reduction of Cx45-puncta in IPL. Our results, combined with those of Maxeiner et al. (2005), suggest that reduced coupling observed in this pathway after Cx36 ko (Güldenagel et al., 2001; Deans et al., 2002) may be attributable not only to loss of Cx36, but also to a small reduction in Cx45 expression or trafficking, which may require Cx45 co-binding with Cx36 and ZO-1 for optimum targeting and/or stability. Because Cx45-containing gap junctions appear to obligately contain Cx36, knock-out of Cx45 from these otherwise bimhomotypic gap junctions also might alter the function and/or stability of the Cx36 that remains in the absence of Cx45.

Fifth, previous studies suggested that most if not all neurons synapsing in the IPL separately express only one or the other connexin isoform (Han and Massey, 2005; Dedek et al., 2006). However, the methods used, including in situ hybridization, single-cell RT-PCR, lacZ reporter expression [by a nonconventional method (Schubert et al., 2005)], or expression of cytoplasmic EGFP as genetic markers (Han and Massey, 2005; Lin et al., 2005; Maxeiner et al., 2005; Dedek et al., 2006), have significant probabilities of both “false-positive” and “false-negative” detection; and as in all techniques, efforts to minimize false positives increase the probability of false negatives by altering the threshold of signal detection. For example, by reducing amplification of lacZ signal so that Cx45-positive cells had from one to a maximum of three lacZ cytoplasmic “dots” (Schubert et al., 2005), weakly expressing cells may have had inadequate amplification to produce a detectable dot, leading to false negatives. Likewise, none of the mRNA-detection methods measure different expression levels that are implied by vastly different sizes (and numbers) of Cx36- versus Cx45/Cx36-containing gap junctions [4–2500 connexons (Kamasawa et al., 2006)]. Our data imply that all amacrine cells, reportedly expressing only Cx36 (Maxeiner et al., 2005), that couple to Cx45/Cx36-coexpressing cells (e.g., cone bipolar cells) must form only Cx36/Cx36 monohomotypic couplings, or that all amacrine cells must also synthesize Cx45 to form bimhomotypic gap junctions, but with Cx45 synthesized at an amount sufficiently low as to have previously gone undetected. Conversely, if all amacrine cells truly lack Cx45, the detection of Cx45-immunofluorescent puncta at apparent cell contacts involving AII amacrine cells must be at tripartite contacts between two other Cx45/Cx36-coexpressing cell processes that are not separately resolvable by light microscopy from the closely associated AII amacrine cell. Such triadic structural arrangements [Kolb (1979), her Fig. 29b] could also account for the reduction in glycine transfer from AII amacrine cells to cone bipolar cells in both Cx36 and Cx45 ko mice (Güldenagel et al., 2001; Maxeiner et al., 2005) if the intermediary neuron expressing Cx36+Cx45 were coupled to glycinergic AII amacrine cells solely by Cx36-containing gap junctions and serially to the nonglycinergic bipolar cells by bimhomotypic Cx45/Cx36 gap junctions. Additionally, this would require that Cx45 ko also disrupt Cx45/Cx36 bimhomotypic couplings.

Overall, our results document that virtually all Cx45-containing gap junctions occur between cell pairs in which both cells coexpress and cotarget Cx45 and Cx36, forming bimhomotypic gap junctions. Thus, interpretation of future coupling experiments involving Cx45-expressing retinal neurons must consider the restricted range of sizes of Cx45-containing gap junctions compared with homotypic Cx36 gap junctions, rather than ascribing observed differences in coupling to heterotypic...
versus homotypic coupling. For example, gating of Cx45 (but not Cx36) is voltage sensitive, and Cx45 has threefold greater conductance (Sohl et al., 2005), potentially allowing differential regulation of gap junction electrical and/or metabolic coupling. Consequently, subcellular regulation of the ratios and abundance of these two connexins in individual gap junctions would allow for finer analog control than available from monohomotypic Cx36 or Cx45 gap junctions, as well as provide a novel mechanism for plasticity of electrical synapses.

References


