Looking Chloride Channels Straight in the Eye: Bestrophins, Lipofuscinosis, and Retinal Degeneration

Recent evidence suggests that Cl\(^{-}\) ion channels are important for retinal integrity. Bestrophin Cl\(^{-}\) channel mutations in humans are genetically linked to a juvenile form of macular degeneration, and disruption of some CIC Cl\(^{-}\) channels in mice leads to retinal degeneration. In both cases, accumulation of lipofuscin pigment is a key feature of the cellular degeneration. Because Cl\(^{-}\) channels regulate the ionic environment inside organelles in the endosomal-lysosomal pathway, retinal degeneration may result from defects in lysosomal trafficking or function. (FIGURE 1). Lipofuscin is a heterogeneous mixture of both proteins and lipids that are partially oxidized (67). It is thought that lipofuscin accumulation is a major causal factor in AMD because it correlates with the severity of photoreceptor degeneration (94) and has been shown to have toxic properties (21, 68). The major fluorescent component of lipofuscin is N-retinylidene-N-retinylethanolamine (A2E), which is a metabolite of retinal visual pigment. A2E has been shown to perturb lysosomal function (21, 68), disrupt membrane integrity by a detergent-like effect (19), and promote programmed cell death (70). Proteomic analysis shows that various components of lipofuscin are derived both from photoreceptors and RPE (67). A possible explanation of this dual cellular origin is provided by the fact that photoreceptor outer segments are renewed by the daily phagocytosis of the tips of photoreceptors by RPE cells (5, 54). It would therefore make sense if lipofuscin accumulation were related to a problem of phagocytosis or lysosomal function (18).

In any case, lipofuscin seems to be a noxious material. Lipofuscin accumulation is also associated with a group of neurodegenerative diseases called neuronal ceroid lipofuscinoses (NCLs). NCLs are lysosomal storage diseases characterized by accumulation of lipofuscin pigment, blindness due to retinal degeneration, psychomotor retardation, and premature death (15, 27, 52). Because some genes associated with NCLs code for proteins critical for lysosomal function (12), this raises the possibility that lysosomal dysfunction may also be involved in AMD.

Best Disease May Be a Model for Macular Degeneration

Both genetic and environmental factors contribute to AMD, but there are a number of early-onset forms of macular degeneration that are exclusively or largely genetic. One of these diseases is Best vitelliform macular dystrophy (also called Best disease), which is inherited in an autosomal-dominant manner and is linked to mutations in the bestrophin-1 gene located on chromosome 11 (11q13). In Best disease, lipofuscin accumulates within macrophages in the subretinal space, in RPE cells, and between RPE cells and photoreceptors. The gene responsible for Best disease was positionally cloned in 1998 from families exhibiting macular degeneration with a juvenile age of onset (48, 58). Over 85 human bestrophin-1 (hBest1) mutations are now associated with macular degeneration (FIGURE 2) (see Ref. 88 and the Human Gene Mutation Database at http://uwmcm1s.uwcm.ac.uk/uwcm/mg/search/133795.html). In addition, inherited abnormal splice variants of hBest1 are associated with autosomal-dominant vitreoretinochorioidopathy, a retinal dystrophy that is associated with defects of ocular...
development such as nanophthalmos (92).

**Best Disease May Be a Chloride Channelopathy**

Although it was initially hypothesized that hBest1 might be a transporter of lipid components of lipofuscin (58), there is now considerable evidence that bestrophins function as Cl⁻ channels. After hBest1 was cloned, Jeremy Nathans’ lab expressed hBest1 in human embryonic kidney 293 cells and showed that its expression was associated with appearance of a novel Ca²⁺-sensitive Cl⁻ current (74). Our lab and Nathans’ lab have now shown that bestrophins from human, mouse, *Xenopus*, *Caenorhabditis elegans*, and *Drosophila* function as Cl⁻ channels when expressed heterologously (62, 74, 79). Subsequently, work from our lab showed that the Ca²⁺ sensitivity was in the physiological range, with an EC₅₀ for Ca²⁺ of ~200 nM (62). The Ca²⁺ sensitivity of hBest1, mouse (m)Best2, and *Xenopus* Best2 are similar, but the Ca²⁺ sensitivity of other bestrophins has not yet been published. Whether bestrophins are regulated by direct Ca²⁺ binding, by calmodulin, or by Ca²⁺-dependent phosphorylation (32) is not known.

**Role of Cl⁻ channels in Best disease**

The suggestion that bestrophins are Cl⁻ channels is particularly intriguing because the hallmark diagnostic feature of Best disease is a decreased slow light peak in the electrooculogram (EOG) (FIGURE 3). The slow light peak is thought to reflect a Cl⁻ conductance in the basolateral membrane of the RPE (24). Because Best1 is localized at the basolateral membrane of the RPE (4, 46), the idea that Best1 is responsible for the basolateral Cl⁻ conductance is particularly attractive.

The mechanisms linking Cl⁻ channel dysfunction to the macular dystrophy that ensues is not clear. This is partly because visual acuity in Best disease patients is highly variable. For example, in one family, an elderly mother...
had normal visual acuity of 20/20, whereas her daughter and three grandchildren had clinical symptoms as early as age 5, even though all had the same genotype (R218C/+) (10). Despite the variable penetrance of disease symptoms, virtually all individuals with bestrophin gene mutations exhibit abnormal EOGs (81). This suggests that other factors, genetic or environmental, must contribute to the development of retinal lesions. However, the fact that an abnormal EOG is a consistent feature of the disease but that degraded visual acuity is more variable provides a strong argument that Cl− channel dysfunction precedes macular degeneration.

Because an abnormal EOG is unambiguously linked to hBest1 mutations, it would be useful to understand the mechanism of the light peak, but this also remains somewhat obscure. To measure the EOG, electrodes are placed on either side of the eye, then the subject is allowed to adapt to darkness and is asked to fixate on one of two light-emitting diodes 30° apart that illuminate alternately at a frequency of ~0.2/s. The difference in voltage measured when the eyes are fixated on the right vs. the left light-emitting diode is thought to represent the voltage across the RPE. During dark adaptation, the potential becomes small (the dark trough) (FIGURE 3). When the ambient lights are turned on, the potential increases slowly to reach a peak in ~5–15 min (light peak). The EOG obviously is an indirect measure of changes in conductance that occur in the retina. The electroretinogram (ERG), which measures the potential across the retina directly, might provide a more direct measurement of the electrical changes. However, there are technical obstacles to using direct-current ERG (dcERG) to record the very-long-duration light peak in humans. Conventional full-field ERGs, which are taken on a much shorter time scale, are often reported to be normal in Best disease. Multifocal ERGs, which use multiple electrodes to sort out spatial differences in retinal function, usually reveal central amplitude

FIGURE 2. Model of hBest1 topology
Hydrophobic segments A–F identified as putative transmembrane domains by the MPEx algorithm are shown. C is not included as a transmembrane domain in this model, because this would place residues 212 and 218 on the wrong side considering the data from TEV protease cleavage. Data are from Ref. 79 and the VMD2 Mutation Database (http://www.uni-wuerzburg.de/humangenetics/vmd2.html).
reductions in patients with clinical symptoms (65). These ERG abnormalities are probably not related to the slow light peak of the EOG but probably reflect degenerative changes secondary to the initial vitelliform lesions.

In animals, dcERG recordings show a slow light peak that has the same time course as the slow light peak in the EOG, suggesting that the slow light peaks in the dcERG and EOG have the same cellular basis. The slow light peak in the dcERG clearly originates from changes in a basolateral conductance in the RPE cells (41). The 5- to 15-min time course of the slow light peak suggests that its mechanisms could be quite complex. It has been proposed that a diffusible substance is released from the neural retina that activates the Cl– conductance in the RPE (41), but the identity of the light substance remains unknown (see discussion in Refs. 24 and 91).

Support for the idea that bestrophin is a basolateral Cl– channel in RPE is provided by recent experiments from Marmorstein's lab (47). In these experiments, an adenovirus encoding either wild-type human Best1 or disease-causing W93C or R218C mutants of hBest1 were injected into the eyes of young rats. The corresponding proteins were found to be expressed in the basolateral membrane of the RPE by immunostaining. Expression of the W93C mutants decreased the light peak of the dcERG without affecting other components of the dcERG. This is the expected result if W93C bestrophin is dominant negative in inhibiting the function of the endogenous wild-type bestrophin. Best disease caused by the W93C mutation is inherited in a dominant fashion, and the W93C mutant has been shown to inhibit wild-type currents in expression systems (62, 74). The effects of the R218C mutant were less than those of the W93C mutant, but the R218C mutation may produce less-severe disease as well. Despite these positive results, the authors express reservations about concluding that bestrophin is the basolateral Cl– conductance, because the effects of overexpression of the W93C mutant on the light peak are considerably less than they expected considering the amount of overexpression. Furthermore, overexpression of wild-type hBest1 did not increase the light peak as expected. However, this might simply mean that the endogenous channels alone produced a maximal response.

**What is the evidence that hBest1 is a Cl– channel?**

As mentioned above, expression of several bestrophins in 293 cells induces a novel Ca2+-activated Cl– current. Unfortunately, identification of Cl– channels by heterologous expression has a troubled history (11, 35). There are a number of proteins, including p64, the CLICs, and the CLCAs, whose Cl– channel function is doubted even though they induce Cl– currents when overexpressed. For the CLICs and p64, skepticism derives from the fact that their structure (small size and only one or no predicted transmembrane domain) seems incompatible with an ion channel. With the CLCAs, there also is disagreement as to whether they are integral membrane proteins (26). A more general problem is that many Cl– currents are biophysically similar and very few specific blockers have been identified. Consequently, it is often difficult to be certain that an expressed protein is not simply upregulating an endogenous Cl– channel: every cell has endogenous Cl– channels and these channels may not be easily separated from the
heterologously overexpressed currents.

With bestrophins, considerable evidence has accumulated to establish that these proteins can function as Cl⁻ channels, although there is the possibility they have other functions as well. Several different arguments support the idea that bestrophins are Cl⁻ channels.

1) Bestrophins from different species expressed in 293 cells produce novel Ca²⁺-activated Cl⁻ currents (62, 74). The type of current that is induced is the same in different cell lines (61). Because one would not necessarily expect upregulation of the same kind of Cl⁻ channel in different cell types, the appearance of similar currents in different cells minimizes (but does not eliminate) the concern over upregulation of endogenous channels (61).

2) In mammals, there are four bestrophin genes (40, 79). In C. elegans, there are at least 25 different bestrophins. Different bestrophins produce currents with different characteristics (74, 79). For example, hBest1 currents have linear current-voltage relationships and are essentially time independent, whereas hBest3 currents strongly inwardly rectify and activate slowly with time (79). These different characteristics provide some confidence that the currents are specific to the type of bestrophin expressed.

3) hBest1 and mBest2 currents are inhibited by the modification of sulphydryl groups with membrane-impermeant MTSET⁺ (61, 63, 79). Modification of ionic currents by MTSET⁺ is a common approach used to identify amino acids that line the permeation pathway (36).

4) Expression of dominant-negative mutants of bestrophin, notably G299E and W93C, inhibit the current induced by expression of the wild-type bestrophin (62, 74), suggesting that the channels are multimers. Semiquantitative co-immunoprecipitation shows that bestrophins form multimeric complexes composed of four or five subunits, as one would expect for a channel protein (74).

5) Point mutations in both hBest1 and mBest2 produce changes in Cl⁻ channel function. Most notably, certain mutations change the selectivity of the channel for anions and/or change channel gating, as described below. Because it is generally agreed that the selectivity of a channel is determined by the channel pore, the ability to change the selectivity by a mutation provides strong evidence that bestrophin is responsible for forming the channel (60).

The Transmembrane Topology of Bestrophin Remains in Question

In discussing the structure-function of bestrophins, it is useful first to consider their transmembrane topology (FIGURE 2). Several different models of bestrophin topology have been proposed based on conventional hydropathy analysis, but the only model to have serious experimental support is the one proposed by Nathans’ lab (79). Nathans’ model for hBest1 proposes four transmembrane domains and a loop. This model is supported by experiments examining the glycosylation of inserted N-glycosylation sites and cleavage of inserted tobacco etch virus (TEV) protease sites. They also inferred the topology by substituting various amino acids with cysteine and testing the effects of the membrane-impermeant sulphydryl reagent MTSET⁺ on the currents (79). Modification of the currents by extracellular MTSET shows that the residue is accessible to the extracellular solution. Although their model is a useful starting point, we feel that the model remains incomplete, because no direct data exists for the sidedness of residues 100–211, but transmembrane prediction by several algorithms, including MPEX (89), suggests that there is a transmembrane domain in this region (FIGURE 2). In their model (79), the segment centered on 138 is not considered to be transmembrane.

Mutagenesis of the second transmembrane domain

We have studied mBest2 as a model for bestrophin Cl⁻ channel function and have mutated a number of residues that either alter the selectivity of the channel for anions or alter channel gating. We have focused on the putative second transmembrane domain (TMD2) partly because this region is one of the most conserved regions in the bestrophins and is the location of a number of disease-causing mutations in hBest1. Also, the finding in hBest1 that residues C69 and N99 are at opposite ends of the predicted transmembrane segment but are both accessible to MTSET applied extracellularly (79) is consistent with TMD2 forming the pore.

Replacing any of several native amino acids in mBest2 TMD2 with cysteine alters the relative permeability of the channel for anions (61, 63). For example, the S79C mutation has a lower relative permeability to SCN⁻ (PSCN⁻/PCl⁻) than wild-type channels, but the relative SCN⁻ conductance (GSCN⁻/GCℓ) is increased. This suggests that the affinity of a binding site in the channel pore is reduced for SCN⁻ relative to Cl⁻ (61). This suggestion is confirmed by measuring the ability of SCN⁻ to compete for Cl⁻ moving through the channel. In wild-type mBest2, SCN⁻ inhibits Cl⁻ currents with an EC₅₀ of 12 mM, whereas in the S79C mutant the EC₅₀ for SCN⁻ is immeasurably large. These data show clearly that bestrophin can function as a Cl⁻ channel, because the binding site for permeant anions is certainly located in the channel pore (60). Substitution of 15 amino acids with cysteine in TMD2 has similar effects to those just described for S79C, but several substitutions in residues in TMD1 (R34), TMD3 (F149, R150), TMD4 (K180, R196, R197, R200), and the loop (D228, W229, I230) produced wild-type currents (Qu and Hartzell, unpublished observations).

Additional evidence that amino acids 72–95 line the pore is provided
by the effect of charged MTS sulfhydryl reagents on cysteine-substituted mBest2 currents. The effects of the positively charged MTSET\(^+\) and negatively charged MTSES\(^-\) on mBest2 currents depend systematically on which residue is changed to cysteine. A helical wheel representation of TMD2 reveals that residues that are insensitive to MTS reagents are clustered on one side of the helix, whereas modifiable residues are concentrated on the other side (FIGURE 4). The fact that changing the electrostatic charge along one side of the TMD2 helix produces changes in Cl\(^-\) conductance provides important evidence that this part of the protein is involved in forming the pore.

Single-channel recordings of bestrophin currents have not yet been published. Unfortunately, this places some limitations on the conclusions that can be made from the mutagenesis experiments. Whole cell conductance is the product of the number of active channels (N), their mean open probability (pₒ), and the single-channel conductance (γ). Although we assume that the effects of MTS reagents reflect changes in γ, another possibility is that these reagents affect channel gating (pₒ). Although we cannot eliminate the possibility that TMD2 is involved in channel gating, the effects of cysteine substitution on anion permeability are more simply explained in terms of channel permeability. Nevertheless, mutations in this domain clearly do alter channel gating, as evidenced by the finding that in the S79T mutation G₅CN/GCI changes with time after switching to SCN\(^-\) (63). Furthermore, we have identified several mutations in TMD2 that dramatically alter channel kinetics (Y. Qu and H. C. Hartzell, unpublished observations).

**Location of mutations in hBest1**

As mentioned above, in mammals there are generally four bestrophin genes (40, 72). In mouse, one of these appears to be a pseudogene (40). The first ~350 amino acids are highly conserved among subtypes and species, whereas the COOH terminus is quite divergent. For example, hBest1 and mBest1 are 83% identical in the first 350 amino acids but only 44% identical in residues from 350 to the end. The high proline content of the COOH terminus (9%) suggests that this region may be important in protein-protein interactions, but specific motif searches have been disappointing.

The mutations in hBest1 that produce Best disease are scattered throughout the first 312 amino acids but are conspicuously absent after amino acid 312 (FIGURE 2). The mutations are concentrated in the NH₂ terminus, TMD2, hydrophobic segment “C” (residues 130–150), the second extracellular loop and the first part of segment “E,” and the ~20 amino acids after the last transmembrane domain. This last region (amino acids 293–311) is highly conserved in all of the bestrophins and is characterized by a cluster of acidic residues. Mutations in 16 of 18 amino acids in this region have been reported to be linked to Best disease. The function of this domain remains unexplored, partly because mutagenesis often produces nonfunctional channels (Qu and Hartzell, unpublished observations). Its acidic nature suggests that it could be a site of Ca\(^2+\) binding, but acidic domains are also a common feature of protein-sorting signals (e.g., see Ref. 43).

**Cl\(^-\) Channels May Play Multiple Roles in Retinal Function**

What are the mechanisms that might explain how defects in bestrophin Cl\(^-\) channel function produce macular degeneration? These mechanisms are summarized in **FIGURE 5** and discussed below.

**Regulation of the fluid surrounding photoreceptors**

RPE cells participate in regulating the volume and ionic composition of the space surrounding photoreceptor outer segments. Epithelial transport across the RPE is mediated by apical (photoreceptor-facing) Na-K-2Cl cotransporters that use the energy from the Na\(^+\) gradient to move Cl\(^-\) uphill into the cell and basolateral Cl\(^-\) channels that allow Cl\(^-\) ions to flow downhill against their electrochemical gradient toward the choroid (6, 23, 64, 66). RPE polarity is opposite to that found in secretory epithelia, such as parotid gland, where the Na-K-2Cl cotransporter and the Na-K-ATPase are located basolaterally and Cl\(^-\) secretion occurs via Cl\(^-\) channels in the apical membrane. Transepithelial movement of Cl\(^-\) is followed by Na\(^+\) as counterion to neutralize charge and water to balance osmotic pressure. Thus there is a net flow of NaCl and water from the neural retina to the choroid (77). Light-dependent changes in photoreceptor ionic currents result in significant accumulation and depletion of ions in the space around the photoreceptor outer segments (34), and it is thought that fluid and ion transport by the RPE helps maintain the composition of this fluid within a range that supports phototransduction. Dysfunction of these transport processes would be expected to have pathological consequences.
membrane, the cytoskeleton and extracellular matrix strengthen the membrane and allow considerably higher gradients to develop. A major mechanism that cells employ to avoid bursting is regulation of their volume.

**Cell volume regulation**

The cell membrane cannot withstand more than 2 kPa osmotic pressure without bursting (28). Although 2 kPa corresponds to ~1 mM osmolyte concentration difference across the membrane, the cytoskeleton and extracellular matrix strengthen the membrane and allow considerably higher gradients to develop. A major mechanism that cells employ to avoid bursting is regulation of their volume in response to osmotic challenge. Cell swelling often activates K⁺ and Cl⁻ channels that result in KCl and water efflux to return the cell to its original volume (regulatory volume decrease) (55). Conversely, cell shrinkage results in regulatory volume increase by mechanisms that involve decreases in ionic conductance.

It is not entirely clear what osmotic forces are experienced by RPE cells in the eye. The volume of the subretinal space increases at light onset as a consequence of K⁺ and water movement out of the RPE (6, 34). Analysis of accumulation and depletion of ions in the subretinal space suggests, however, that the change in osmolarity may be small (16). Nevertheless, RPE cells express volume-regulated anion channels (9, 38) and the currents induced by bestrophins expressed in 293 cells are very sensitive to changes in cell volume (22). Another possible role for volume-sensitive Cl⁻ channels is in regulation of cell volume as a consequence of ingestion of photoreceptor discs as discussed below.

**Intracellular Cl⁻ channels**

Although it is clear that bestrophins can function as Cl⁻ channels at the plasma membrane, is it possible that they also have an intracellular function? Several of the CIC Cl⁻ channels, such as CIC-3 and CIC-7, have both plasma membrane and intracellular functions (35). One function of intracellular Cl⁻ channels is to regulate the pH of intracellular membrane compartments. For example, the luminal pH of the membrane compartments decreases along the endosomal pathway. The acidic luminal environment develops as a result of the vesicular proton pump that pumps protons against their concentration gradient into the organelle. To acidify the lumen, however, a counterion shunt pathway is required to dissipate the transmembrane potential that develops as protons accumulate in the lumen. This counterion shunt appears to be provided largely by Cl⁻ channels, at least in the endosomal pathway (20). The luminal pH of the vesicle is important in membrane trafficking.

**FIGURE 5. Functions of chloride channels in RPE**

Depending on the conditions, retinal pigment epithelium (RPE) cells can function as an absorptive or secretory epithelium. The basolateral Cl⁻ channel responsible for the light peak regulates membrane potential and depolarizes the RPE in light. Intracellular Cl⁻ channels regulate vesicular pH and intraluminal Cl concentration, which affect both enzymes and receptors for cytoplasmic signaling molecules. CSP, Cl⁻-sensitive protein; PSP, pH-sensitive protein. Although there is no compelling evidence for Cl⁻ channels in the apical membrane, we hypothesize that Cl⁻ channels in the apical membrane may be involved in regulating the ionic composition of the solution bathing the photoreceptors under certain conditions. Volume-regulated anion channels (VRAC) regulate the cell volume. VRAC is presumably located in the basolateral membrane. Ca²⁺ may regulate the basolateral Cl⁻ channel, VRAC, and intracellular Cl⁻ channels as discussed in the text. The intracellular Cl⁻ channel is shown as a H⁺/Cl⁻ exchanger because recent evidence has shown that certain intracellular CIC channels are transporters, as discussed in the text.
Cl– channels are likely to contribute not only to the maturation of the phagolysosome by controlling its pH and water content (1, 42) but also in regulating membrane traffic along the endosomal-lysosomal pathway. In liver cells, phagocytosis is accompanied by large osmolyte fluxes via anion channels and transporters (87), and changes in extracellular osmolarity alter phagocytosis (82). The role of Cl– channels in phagocytosis is underscored by the observation that phagocytosis is blocked by the Cl– channel blocker tamoxifen (44).

Another emerging function of intracellular Cl– channels is regulation of Ca2+ movement across the endoplasmic reticulum by providing a counterion pathway. Luminal Ca2+ concentration is important for protein folding and quality control in the endoplasmic reticulum (51), and Ca2+ imbalance in the endoplasmic reticulum can lead to cell death (80). Cl– channels have been shown to be present in sarcoplasmic reticulum, and Cl– channel blockers alter Ca2+ uptake into the sarcoplasmic reticulum (59). One would speculate that if Ca2+-activated Cl– channels are present in the endoplasmic reticulum, increases in cytosolic Ca2+ would activate these and facilitate Ca2+ flux across the endoplasmic reticulum membrane.

Significance of calcium regulation

Overexpressed bestrophins have an EC50 for Ca2+ in the range of 200 nM (61, 62). If this represents the Ca2+ affinity of native channels, bestrophins would be expected to be at least partially activated at typical resting Ca2+ concentrations of 50–200 nM. At present, the physiological significance of the Ca2+ sensitivity of bestrophins is speculative. Although there are Ca2+-stimulated Cl– conductances in RPE (see references in Ref. 33), there is no direct evidence that the light peak of the dcERG is Ca2+ dependent. The role of Ca2+ in cell volume regulation is equivocal, although there is good evidence that Ca2+ may facilitate the regulatory volume decrease but is not obligatory (2). Finally, there is no information about how cytosolic Ca2+ might regulate Cl– channels in intracellular vesicles.

Can Best Disease Provide Insights into the Mechanisms of AMD?

Although lipofuscinosis is thought to be a causal factor in AMD, it is clear that the consequences of lipofuscin accumulation differ in Best disease and AMD (e.g., Refs. 25 and 56). AMD is typically characterized by deposits of debris called drusen between the RPE and Bruch’s membrane (7, 25). Drusen contains proteins characteristic of immune processes, suggesting that it is produced as a consequence of inflammation (3, 13, 29, 30). This is interesting in light of the fact that specific polymorphisms in complement factor H correlate strongly with susceptibility to AMD (17, 30, 31). In contrast, in Best disease there are few or no drusen. If lipofuscin accumulation is causal to drusen formation in AMD, why is there only modest drusen formation in Best disease? Does it reflect complement factor H genotype, for example? The answer may also be hidden in our limited understanding of lipofuscin. The composition of lipofuscin appears to be different in AMD and Best disease (45), and its distribution is different. In AMD, the vitelliform egg-yolk-like lesions that are characteristic of Best disease are usually not present (56, 84). However, vitelliform lesions have been reported in individuals who have no known hBest1 mutations (65). Despite these differences between Best disease and AMD, elucidating the mechanisms of lipofuscin accumulation and toxicity will be an important step in understanding these diseases.

dysfunction of some CIC Cl– Channels Produces Retinal Degeneration

There seems to be an interesting relationship between Cl– channels, lipofuscinosis, and retinal degeneration. RPE cells express a variety of different kinds of Cl– channels in addition to bestrophins. CIC-2, CIC-3, CIC-5, and CFTR have been identified by RT-PCR and Western blotting (86, 90). Electrophysiologically, currents resembling CIC-2 and CFTR, as well as Ca2+-activated and volume-activated Cl– currents, have been described (24, 33). Interestingly, in mice that have had the CIC-2, CIC-3, or CIC-7
Cl– channels knocked out, the retina degenerates (8, 39, 71). Retinal degeneration in the CIC-2 knockout may be due to an RPE dysfunction because RPE short-circuit current is reduced in CIC-2 knockout mice (8). The mechanisms by which knockout of these genes produce retinal degeneration remain speculative. CIC-2 and CIC-3 are located both on the plasma membrane and in intracellular compartments. Their plasma membrane roles certainly include transepithelial transport. Dysfunction of these channels, therefore, would likely result in abnormal composition of fluid surrounding photoreceptors.

In addition to retinal degeneration, the CIC-3 and CIC-7 knockouts have a more generalized neurodegenerative disease resembling NCL (15, 27, 37, 52, 93). NCLs have been linked to mutations in eight genes (CLN1–CLN8). Several of these have been identified as lysosomal enzymes or components of lysosomes. CIC-7 is also localized in lysosomes (37). One hypothesis for the mechanism by which CIC-7 deletion produces an NCL-like disease and retinal degeneration is related to the role of Cl– channels in regulating the pH of intracellular compartments (20). This hypothesis is attractive because changes in lysosomal pH have long been known to be associated with retinal degeneration. An increase in lysosomal pH can be produced by chloroquine, which is a weak base that accumulates in lysosomes. A common side effect of chloroquine therapy for autoimmune diseases is retinopathy associated with accumulation of lipofuscin (69, 75, 76).

Surprisingly, however, changes in lysosomal pH do not explain the NCL phenotype of the CIC-7 knockout, because lysosomal pH is normal in these animals (37). An alternative possibility that should be considered is that Cl– concentration in the vesicle might itself be an important regulator of vesicular function, as we proposed previously (20). In yeast, for example, a CIC-like Cl– channel is implicated in regulating the Cl– concentration of Golgi compartments that is required for Cu2+ loading of an oxidase enzyme (14). (See Notes added in proof, below).

Although the CIC-3 deletion was postulated to produce lipofuscinosis by reducing lysosomal enzyme activity or blocking maturation of lysosomes as a result of an elevated organellar pH (93), CIC-3 is not concentrated in lysosomes like CIC-7 but rather is found in endosomes and synaptic vesicles. This suggests that the phenotype of the CIC-3 knockout may be caused by another mechanism. Furthermore, another mouse strain with a similar CIC-3 deletion does not exhibit an NCL phenotype (37), implying that genetic background or other factors may be important.

Conclusions

The relationship between various chloride channels and lipofuscinosis suggests that neural degeneration in NCL and photoreceptor degeneration in macular degeneration may involve similar mechanisms. Our working hypothesis is that Cl– channels may have multiple functions, both at the plasma membrane and in intracellular organelles. Defects in Cl– channel function could result in retinal degeneration as a consequence of problems with cell volume regulation, function and trafficking of intracellular organelles, or regulation of the extracellular fluid surrounding outer segments. However, the fact that lipofuscin accumulation occurs in NCL as a consequence of lysosomal dysfunction suggests that the retinal degeneration that occurs in Best disease is also a lysosomal problem, especially because RPE cells are such active phagocytes. Understanding the mechanistic relationship between lipofuscin accumulation and chloride channels will provide valuable insights into the process of AMD (see below).

Notes added in proof

Recent data that certain intracellular CICs are proton-Cl– exchangers raises interesting questions about the role of CICs in regulating vesicular pH. (58a, 66a).

An excellent review on the physiolo-

gy of RPE cells has recently been published. Interested readers should consult the review for more detail on this interesting cell type (73a).

We would like to thank Drs. Jeremy Nathans, Machelle Pardue, and Neal Peachey for comments on the manuscript.

The research in our lab is supported by National Institutes of Health Grants EY-14852 and GM-60448 and by the American Health Assistance Foundation.

References


66a. Scheel O, Zdebik A, Lourdel S, and Jentsch TJ. Physiology


