Common Cell Biologic and Biochemical Changes in Aging and Age-Related Diseases of the Eye: Toward New Therapeutic Approaches to Age-Related Ocular Diseases

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Reviews of information about AMD, cataract, and glaucoma make it apparent that while each eye tissue has its own characteristic metabolism, structure, and function, there are common perturbations to homeostasis that are associated with age-related dysfunction. The commonalities appeared to be biochemical stresses and their sequelae. Recognition of shared etiologic factors for age-related deilities allows rationalization of comparable risk factor-disease incidence relationships—such as nutritional risk factors for AMD and cataract (as well as cardiovascular disease and diabetes)—and informs about potential new therapeutic avenues, such as stress reducers (i.e., antioxidants) and/or proteolysis enhancers. It also maximizes the return on the investment in research effort and costs. For example, drugs or nutrients that protect against AMD may also prove effective against cataract, glaucoma, or/and other age-related neurodegenerative deilities.

This article summarizes cell biologic and biochemical changes in aging and age-related diseases of the eye. Clearly, this is a larger challenge with a richer literature than can be properly treated in a short review such as this. In this short review, we focus on age-related stresses and current and anticipated means to diminish the stress. Recognizing that almost all age-related diseases such as Alzheimer and Parkinson diseases, cataract, AMD, glaucoma, diabetes, and the premature aging diseases such as progeria, have in common the accumulation of damaged proteins, we select three aspects of age-related biochemical changes that are common to most eye tissues: oxidative stresses; problems associated with and/or due to damaged proteins that accumulate in the retina, lens, and cornea; and intracellular degradative capacities that usually keep levels of damaged proteins in check in early life or when tissues are not stressed, but that may fail upon stress or aging (Figs. 1, 2). We offer apologies to investigators whose work we do not cite or can acknowledge only via reviews.1

The most rapidly growing segment of many societies is the elderly. The prevalence of cataract, AMD, and glaucoma accelerates with age. Among those who are aged 75 years or older, prevalence rates of cataract, AMD, and glaucoma are approximately 60%, 15%, and 20% of the population, respectively. These estimates almost double for people aged just 10 years older. Like most tissues in general, most eye tissues suffer from the accumulation of damaged proteins. Such accumulation appears to involve post-synthetic modifications to proteins and limits on the proteolytic capacities that are normally available to degrade and remove the altered or obsolete proteins before they transform into cytotoxic aggregates. Collectively, we call the sum of synthesis, post-synthetic modification, editing and removal of proteins “proteopoiso.” Compromises to proteopoiso are also thought to be etiologic for many age-related neuropathies and premature aging syndromes.1–7 Herein, we work our way from the anterior of the eye, cornea, through to the lens and on to the posterior segment or retina, recalling common themes of age-related changes and protein quality control.

AGE-RELATED CHANGES IN THE CORNEA, LENS, AND RETINA

The cornea is a multilayered tissue containing three distinct cellular layers, epithelium, stroma, and endothelium, and two membrane structures: Bowman’s layer, separating the epithelium and stroma; and Descemet’s membrane, separating the stroma from the endothelium. The major functions of the cornea are to protect the rest of the eye from environmental insults and to refract light.

Structural and biochemical changes have been noted in all layers of the cornea upon aging. The corneal epithelium becomes more permeable with age,8 possibly due to alterations in the distribution of α6 and β4 integrins, transmembrane receptors that mediate the attachment between a cell and its surroundings.9 Age-related alterations in the human (diurnal) cornea appear to involve cumulative, prolonged ultraviolet radiation exposure as well as stresses that are associated with aging per se. This leads to the generation of reactive oxygen species that, in turn, cause oxidative stress. Accordingly, it is not surprising that protein oxidation is a frequent insult to the cornea.10 This involves advanced glycation-end products.


(AGEs) that form due to a nonenzymatic reaction between proteins and aldehydes and ketones, most of which are derived from sugars. Levels of AGEs increase upon aging in corneal collagen, lens, and probably all eye tissues and may be further increased by diabetes or due to consuming high-glycemic index diets. AGE-modified collagen may contribute to the increase in collagen fibrils and decreased corneal flexibility observed upon aging. Age-related thickening of Bowman’s layer and Descemet’s membrane also involves post-synthetic modification.

Additional evidence of age-related oxidative damage derives from analyses of genetic material. Genomic and mitochondrial DNA are damaged with age and corneas from older donors show increased 8-OHdG, a marker of DNA oxidation, a consequence—at least in part—of the age-related compromise in DNA damage repair capacity.

Ascorbate and glutathione are important nonenzymatic antioxidants in the cornea. Ascorbate levels are significantly higher in the cornea than in the serum or aqueous humor. Surprisingly, there is a trend toward increased ascorbate, glutathione peroxidase, and the antioxidant cytoglobin in aged human corneas. However, there was a decrease in both mRNA and protein expression of superoxide dismutase-1 and γ-glutamyl–transpeptidase activity. Similarly, in aged rabbit corneas, the levels of glutathione peroxidase, superoxide dismutase, and catalase were significantly decreased. Because these antioxidant enzymes provide critical protection against oxidative stress, age-related losses in their activity would confer enhanced susceptibility to stress.

With mounting stress, it is not surprising that damaged proteins and other potentially harmful moieties also accumulate. Mutations also cause accumulation of abnormal proteins, many of which have been etiologically associated with disease. An example is optineurin. Although incompletely characterized at present, optineurin appears to have roles in apoptosis, inflammation, vasoconstriction, morphogenesis, membrane, and vesicle trafficking, as well as in transcription activation. Mutations in optineurin have been related to risk for glaucoma, and are also found in inclusions in patients with amyotrophic...
lateral sclerosis (ALS). The accumulation of altered proteins is thought to be exacerbated by insufficient proteolytic or other degradative capacity and etiologically related to many premature age-related diseases including Parkinson, Alzheimer, and ALS. 

Corneal epithelial cells show increased expression of p21WAF1 and p16INK4A and senescence-associated beta galactosidase during aging, consistent with diminished proliferative capacity and cell density with age. Since p21WAF1 is a substrate of the ubiquitin proteasome proteolytic pathway, it is possible that intracellular proteolytic capacities are compromised upon aging (see Age-Related Changes In Proteolytic Capacities During Aging In Cornea, Lens And Retina section).

Light coming from the cornea must pass through the fiber cells of the lens nucleus en route to the retina. Of all the tissues in the eye, it is probably easiest to recognize deficits in proteopoen in the lens. Fiber cells are functionally analogous to a fiber optic. When young, they are filled with a clear solution of native proteins. The lens is also equipped with very high levels of glutathione, ascorbate, and antioxidan enzymes. However, upon aging and stress, these levels decline and the antioxidant enzymes are rendered less active. Consequently, proteins are gradually modified, often by oxidation, deamidations, racemizations, and they lyse or aggregate and precipitate. In recent years, there has been increased interest in the damage that is caused by elevated levels of dietary sugars, or AGEs because elevations in intake of carbohydrates has recently been related to enhanced risk for cataract, AMD, cardiovascular disease, and diabetes. The rates of accumulation of many of these post-synthetic alterations appear to accelerate upon aging (Figs. 1, 2).

The retina is composed of a myriad of cell types. They can be very roughly divided into neural retina, RPE, and the choroidal vessels that feed the rear of the retina. The choroidal vessels at the rear of the eye supply nutrients and oxygen to the outer layer of the retina, and actively transport waste away from the retina. Clearly, they are essential for maintaining retinal health, but upon aging, they are partially lost in humans. In contrast with thinning in humans, in mice there is evidence of increased choroidal thickness upon aging. Retinal pigmented epithelial cells and photoreceptors are also lost, particularly in AMD.

Multiple studies have identified oxidative stress as an etiologic factor in AMD. The retina a fertile environment for oxidative stress. This is due to the presence of two blood supplies, the highly oxygenated environment, along with the presence of high levels of photosensitizers and readily oxidizable lipid, protein and carbohydrate substrates. This is exacerbated by a huge proteolytic burden, particularly in the RPE, due to the requirement to degrade the tips of photoreceptor outer segments that are shed nightly. Oxidative stress is indicated by the contents of basal laminar deposits and drusen that herald the onset of AMD and by the marked increase in risk for AMD in smokers. Additionally, Handa and colleagues found evidence of AGE modification of choroidal proteins in an aged donor, who exhibited no age related eye disease, and this was expanded by observations of elevated levels of AGEs and harbingers of AMD in older animals that consumed higher glyceric index diets. Importantly, we find a systemic burden indicated by the higher levels of AGEs throughout the eye and many bodily tissues of mice that consumed higher GI diets. Emphasizing that this is a diet GI-AMD risk relationship, there is increased risk for each category of AMD in people who consume the highest GI diets. Since people who consume higher GI diets are at increased risk of AMD, as well as cardiovascular disease and diabetes, it is likely that the accumulation of AGEs and disease are mechanistically linked and that treatments for one malady may bear benefits for other etiologically linked debilitating. Importantly, findings from the Age-Related Eye Diseases Study 2 indicate that intake of elevated levels of vitamins C and E as well as zinc and lutein confer some protection against progress of intermediate to advanced AMD. Clearly, it would be of greatest interest to find the means to avoid onset of AMD.

Genetic variations in complement factor H confer major risk for AMD, but the mechanism is unknown. The hypothesis that oxidative stress or its sequelae are involved in risk for AMD was corroborated by Weisemann’s recent observation that mutant complement factor H cannot detoxify lipid oxidation products as effectively as the normal protein. This is the first mechanistic link between robust epidemiologic associations regarding risk for AMD and the multiple studies that have associated genetic mutations, particularly genes that regulate immune and inflammatory responses, with risk for AMD. The relationship between inflammation and risk for AMD is also supported by observations of increased numbers of macrophages in the choroid of aged mice, as well as increased levels of prostaglandin, PGE2, and its receptor PGE2-EP2 in the choroid of aged rats. The age-altered cytokine profiles and macrophage responses to retinal laser insult inform about new targets for therapy.

AGE-RELATED CHANGES IN PROTEOLYTIC CAPACITIES DURING AGING IN THE CORNEA, LENS, AND RETINA

In addition to post-synthetic modifications, damaged or obsolete proteins may accumulate because they are not recognized as obsolete, or because of insufficient proteolytic capacity. Such proteins may aggregate, as is observed in cataract, or be otherwise cytotoxic. Recognition of damaged proteins in the cytoplasm is usually accomplished by ubiquitination. In the ubiquitin pathway, ubiquitin is attached to substrates by the sequential activities of E1 (ubiquitin-activating enzymes); E2s (ubiquitin-conjugating enzymes); and E3s (ubiquitin ligases). The exquisite selectivity of ubiquitin conjugation is achieved by the combinatorial activities of dozens of E2s together with hundreds of E3s. Ubiquitinated proteins are delivered to the proteasome where the ubiquitin is removed and recycled and the substrate is degraded to peptides that are eventually reduced to amino acids by aminopeptidases. Selectivity is also achieved by the many deubiquitinating enzymes that, by regulating the extent of ubiquitination, control access of the ubiquitinated substrate to the proteasome. An important observation regarding links between proteolytic capacities and oxidative status derives from observation that all the conjugating enzymes of the UPS are sulfhydryl enzymes and that the GSH/GSSG ratio controls their activities.

A parallel—if less selective—pathway involves autophagy. In this pathway, parts of the cytoplasm are engulfed by a membrane and delivered to the lysosome for degradation. Multiple recent papers document that the UPS and autophagic pathways work in concert. Calpains also complement the UPS and autophagosomal pathways and recent discoveries indicate that these pathways may be linked (Liu and Taylor, unpublished observations, 2015).

What are the proteolytic capacities in cornea, lens, and retina and what is their fate and functional status during aging and upon stress? In the cornea, the larger literature to date deals with extracellular proteases and collagenolysis. Evidence regarding additional proteolytic capacities is being developed, in part, via high-throughput analytical methods.

Since protein aggregation and precipitation is clearly related to cataract, much effort has been devoted to documenting
proteolytic capacities in the lens and their fate upon oxidative stress and aging. While lenses contain all of the ubiquitin, lysosomal, and calpain machineries in the young nucleated cells, by virtue of the lens fibers degrading their organelles, any opportunity for renewal and much of the autophagic lysosomal proteolytic capacity is lost during differentiation and aging. There are also some species differences, such as in the complement of calpains in lenses from different species. Ubiquitin-conjugating activity decreases in old rat lens as compared with younger rat lens. The conjugating activity is also decreased from the outer cortex (newer fiber cells) to the lens nucleus (oldest fiber cells). In the human lens, the three peptidase activities of the proteasome did not change with aging, but they are found at decreased levels in cataractous lenses, consistent with a requirement for these activities to rid lens cells of damaged proteins. As noted above, even when the cells have the full complement of ubiquitination enzymes, increased oxidative stress can diminish their efficacy (Fig. 1). Similarly, oxidative modifications to substrates may alter access to these systems. Thus, it has been demonstrated that while mild oxidative stress or deamidation makes some substrates better proteasomal substrates, glycation demonstrated that while mild oxidative stress or deamidation substrates may alter access to these systems. Thus, it has been observed that chaperones clearly interact with the cellular proteolytic machineries. There appears to be an age-dependent decline in chaperone-mediated autophagy, at least in liver. This seems to be due to a decrease in the lysosomal receptor Lamp2A. Importantly, restoration of chaperone-mediated autophagy in aging liver improves cellular maintenance and hepatic function, including the removal of modified proteins. Evidence for salutary effects of chaperones is accumulating in eye tissues.

**Unmet Needs and Opportunities**

Taken together, it is clear that a vast array of antioxidant and proteolytic capacities maintain homeostasis, when they are functional and unimpaired. However, upon aging or stress, these capacities are inactivated and become insufficient. Thus, there ensue a vicious cycle of stress, accumulation of damaged proteins, and diminished proteolytic capacity that leads to accelerated accumulation of cytotoxic materials, and disease. We know that adequate nutrition is crucial for maintenance of eye function. Accordingly, massive educational campaigns should be instituted that relate risk for blindness to nutrition, smoking, obesity and exercise. These are bound to bear positive results. We could eliminate much age-related disease if salutary behavior changes or treatment could be implemented earlier. To encourage this, we need to determine biomarkers that will allow us to anticipate incipient disease. Current technologies make this an achievable objective. Education about healthy lifestyles and eating habits is probably the least costly and most effective investment with regard to achieving prolonged eye function. It has recently been demonstrated that electron recycling agents can delay mitochondrial disease. Since the age-related debilities noted here all have origins in oxidative stress, it is likely that such agents can also delay formation of the offending moieties and/or preserve function of the degradative machines (UPS, autophagic/lysosomal) that remove oxidized products. Recent findings that degradation of neurodegeneration-related proteins can be enhanced should be translated to therapy for other age-related protein precipitation diseases including cataract, AMD, and possibly glaucoma. New and more potent antioxidants can be discovered or constructed for delivery to eyes to enhance protection against oxidative stress. Elucidating components of age-related vision loss that are not due to retina or lens damage will provide needed understanding and open up new alternatives for vision restoration for people with neurologic damage.

Given the promise for success and the low cost of this research relative to the gains in sight maintenance from its successful completion, it is clear that additional funding would be a wise investment.

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**References**


