In the vertebrate eye, limiting oxidation of proteins and lipids is key to maintaining lens function and avoiding cataract formation. A study by Serebryany et al. identifies a surprising contributor to the eye’s oxidative defense in their demonstration that γD-crystallin (HyD) functions as an oxidoreductase and uses disulfide exchange to initiate aggregation of mutant crystallins that mimic oxidative damage. These insights suggest a mechanism by which a dynamic pool of closely packed proteins might avoid oxidation-driven protein-folding traps, providing new avenues to understand the basis of a human disease with global impact.

The eye lens walks on the wild side of life. Lens proteins (1) do not turn over as in other cells, and, indeed, carbon-dating of eye proteins is often used to determine the age of a particular animal. One consequence of this literal lifetime is that any damage that occurs accumulates over a multiyear timeline. To avoid damage, the lens has an impressive oxidation defense system, including the highest concentrations of glutathione (GSH) found anywhere in the body, as well as biosynthetic and regenerative systems to maintain this high GSH concentration (2). However, as we age, a chemical barrier develops to prevent GSH diffusion from the cortex (outside) into the nucleus (center), isolating the oldest, and arguably most vulnerable, cells in the nucleus of the lens (see Ref. 2 and references therein). As a result, the effectiveness of this defense system erodes (2) as GSH breaks down (3), leading to increases of oxidized thiols in the form of disulfides, both mixed (3) and intermolecular (see Ref. 2 and references therein), novel post-translational modifications, and cross-links based upon cysteine oxidation (4). This eventually leads to the loss of lens function due to protein aggregation and cataract formation, which remains the major cause of blindness worldwide (http://www.who.int/blindness/causes/priority/en/index1.html, accessed September 6, 2018).

Why would the lens evolve a built-in GSH barrier designed to activate later in life if it’s a major redox buffer? Is this the smoking gun for the involvement of other as yet unknown players in lenticular redox homeostasis? An answer to this question has been provided by Serebryany and colleagues (5), who have previously studied the crystallins. These are major components of the lens proteome and present in the lens at concentrations of up to 600 mg/ml in order to deliver the graded refractive index needed to focus light onto the retina. Additionally, oxidative damage to the crystallins in the form of increasing disulfide bonds is directly correlated with cataracts. The β- and γ-crystallins, two of the three major crystallin classes, belong to an ancient superfamily (6) characterized by Greek key motifs and calcium- and transition metal–binding properties and are typically thought of in terms of their structure and refractive properties. Specifically, the close packing between these proteins and their very high concentration (see Ref. 6 and references therein) underpins lens function but runs the risk of initiating aggregation. In their previous work, Serebryany and colleagues (7) explored the aggregation of the W42Q mutant of HyD, which mimics a cataract-associated mutation. They discovered that the WT isomorf of HyD could promote aggregation of the W42Q mutant in a disulfide bond–dependent manner. In their new study, Serebryany et al. (5) now present evidence that this function involves protein disulfide exchange, defining an unexpected oxidoreductase activity of HyD.

HyD contains six cysteine residues, with the cysteine pair at positions 108 and 110 resembling the “CXC” motif of some thioredoxins and disulfide isomerases, as noted by Serebryany et al. (5). The authors confirmed these residues do form a disulfide bond upon oxidation and that this event is required to promote aggregation of the W42Q mutant. Furthermore, the authors explicitly demonstrated disulfide exchange between HyD and the W42Q mutant, although the newly formed disulfide occurs between Cys^{32} and Cys^{41} of the W42Q mutant. Interestingly, the same disulfide transfer does not occur between WT HyD molecules.

It is an open question as to what triggers the disulfide transfer. The authors demonstrate that the Cys^{108}–Cys^{110} bond destabilizes the WT protein, so relief from this conformational strain may facilitate the reaction, but that does not explain why transfer only occurs to the mutant sequence. One possibility is that the Trp to Gln mutation adjacent to Cys^{41} lowers the redox potential of the Cys^{32}–Cys^{41} disulfide, driving electron flow from the Cys^{32},Cys^{41} pair to the Cys^{108}–Cys^{110} bond (Fig. 1). The authors also suggest conformational changes in the mutant could facilitate disulfide transfer and additionally promote...
aggregation of oxidized W42Q but not WT H\(\text{yD}\). The impact of protein concentration has not been considered as the experiments were conducted at protein concentrations two orders of magnitude less than in the eye lens. Nevertheless, in their model, the authors suggest that a “hot potato” competition is in play with the oxidation of Cys\(^{108}\)–Cys\(^{100}\) leading to a less stable W42Q H\(\text{yD}\) and predisposing it to aggregation (7). The model also implies that WT H\(\text{yD}\) can act like a catalyst, converting many molecules of mutant H\(\text{yD}\). Prevention of H\(\text{yD}\) oxidation might block aggregation of the W42Q mutant. What, however, should we make of these results in the absence of mutations or pre-existing damage?

To us, these data lend themselves to a larger scale “hot potato” model in which \(\beta\)- and \(\gamma\)-crystallins are continually rubbing shoulders within a high-protein concentration, close-packed, disulfide-sharing, dynamic protein network. It is akin to the annual emperor penguin huddle in Antarctica, but on a completely different length and temperature scale! The oxidoreductase function provides the capability to resolve undeclared contacts, i.e. disulfide bonds, by sharing them within the crystallin protein network as though they are at a never-ending “meet and greet” party. Deleterious electrons are either removed by appropriate sinks elsewhere or return so slowly that they do not interfere with normal function. As humans (and mice) age, however, disulfide formation shifts from intra- to intermolecular configurations, a trend that is strongly linked to cataractogenesis (2). For example, an experiment meant to mimic aging demonstrated that in an oxidized H\(\text{yD}\) containing an N-terminal methionine, Cys\(^{111}\) (equivalent to Cys\(^{110}\)) forms an intermolecular disulfide with either Cys\(^{111}\) or Cys\(^{19}\) (i.e. Cys\(^{18}\)) in an adjacent H\(\text{yD}\) chain (8). In these cases, the “hot potato” cannot be passed further and aggregation occurs.

It is possible that other \(\gamma\)-crystallins could have similar oxidoreductase-like properties as H\(\text{yD}\), building the possible avenues by which disulfides could be dispersed. For example, although it is not expressed at the same high levels as H\(\text{yD}\), \(\gamma\)\(\alpha\)-crystallin has the same cysteine arrangement. In human \(\gamma\)S-crystallin (H\(\text{yS}\)), there is an N-terminal \(\beta\)-strand DCDCDC (Cys\(^{23,25,27}\)) sequence. H\(\text{yS}\) also has the potential to form intermolecular disulfides as oxidation of Cys\(^{83}\) in H\(\text{yS}\) along with one of these N-terminal cysteines has been shown to encourage its dimerization with \(\beta\)-crystallins due to a conserved disulfide interface (9). It will be fascinating to learn more about these proteins and, through them, aging compensation mechanisms more generally.

In conclusion, the Serebryany et al. (5) paper demonstrates an exciting new function to add to the repertoire for the crystallin family. It emphasizes further the importance of oxidative mechanisms in lens cataractogenesis. The article also provides new impetus to explore oxidation chemistry–based solutions to the global challenge of treating cataracts, particularly in parts of the world where advanced surgery and laser treatments are unavailable, but also where limited medical resources could be deployed more efficiently in favor of pharmacological solutions. Indeed, recent reports have shown the efficacy of redox-active lipoic acid to prevent presbyopia (10) and an antioxidant mixture to ameliorate cataractogenesis in animal models (11), providing new opportunities to augment the redox balance in the eye lens.

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Editors’ Pick Highlight: Redox–detox in the lens

Figure 1. Model for disulfide transfer from oxidized H\(\text{yD}\) to reduced W42Q H\(\text{yD}\). The Cys\(^{108}\)–Cys\(^{100}\) and Cys\(^{32}\)–Cys\(^{41}\) potential disulfides in H\(\text{yD}\) are shown. The Trp to Gln mutation adjacent to Cys\(^{31}\) is shown in red.