INTRODUCTION

Although vitreous is the largest structure within the eye, constituting 80% of the ocular volume, investigators of vitreous anatomy are hampered by two fundamental difficulties:

- Any attempts to define vitreous morphology are efforts to “visualize” a tissue that is invisible by design (Fig. 6-46-1).
- The various techniques that were employed previously to define vitreous structure were flawed by artifacts induced by tissue fixatives, which caused precipitation of hyaluronan (formerly called hyaluronic acid), a glycosaminoglycan.

The development of slit-lamp biomicroscopy by Gullstrand in 1912 was expected to enable clinical investigation of vitreous structure without the introduction of the aforementioned artifacts. Yet, a widely disparate set of descriptions resulted because of the first of the inherent difficulties just described; that is, the vitreous is largely invisible. This problem even persists in so-called modern investigations. Consider, for example, that in the 1970s Eisner1 described “membranelles” and Worst2 “cisterns,” in the 1980s Sebag and Balazs3, 4 identified “fibers,” and in the 1990s Kishi and Shimizu5 found “pockets” in the vitreous. The discrepant observations of the last-mentioned group have now been explained largely as an age-related phenomenon with no relevance to the inherent macromolecular structure or anatomy.6, 7

MOLECULAR MORPHOLOGY

Supramolecular Organization

Vitreous is composed of a dilute meshwork of collagen fibrils [Fig. 6-46-2] with interspersed extensive arrays of long hyaluronan molecules.8–10 The collagen fibrils provide a solid structure that is “inflated” by the hydrophilic hyaluronan.11 Rheological observations also suggest the existence of an important interaction between hyaluronan and collagen.12 Balazs has hypothesized that the hydroxylysine amino acids of
collagen mediate polysaccharide binding to the collagen chain through O-glycosidic linkages. In cartilage, “link glycoproteins” have been identified that interact with proteoglycans and hyaluronan. Supramolecular complexes of these glycoproteins are believed to occupy the interfibrillar spaces. Bishop has elegantly described the potential roles of type IX collagen chondroitin sulfate chains, hyaluronan, and opticin in the short-range spacing of collagen fibrils and how these mechanisms might break down in aging and disease. Many investigators believed that hyaluronan–collagen interaction occurs on a “physicochemical” rather than a “chemical” level. Reversible complexes of an electrostatic nature between solubilized collagen and various glycosaminoglycans could indeed form, since electrostatic binding between negatively charged hyaluronan and positively charged collagen could occur in vitreous.

**VITREOUS ANATOMY**

**Macroscopic Morphology**

In an emmetropic adult human eye the vitreous is approximately 16.5 mm in axial length with a depression anteriorly just behind the lens (patellar fossa). The hyaloidecapsular ligament of Weiger is the annular region (1.2 mm in width and 8–9 mm in diameter) where the vitreous is attached to the posterior aspect of the lens. Erggelet’s or Berger’s space is at the center of the hyaloidecapsular ligament. The canal of Cloquet arises from this space and courses posteriorly through the central vitreous at the center of the hyaloideocapsular ligament. The canal of Cloquet attaches to the posterior aspect of the lens. Erggelet’s or Berger’s space (retrolental space of Erggelet) is the former site of the hyaloid artery in the embryonic vitreous. The former lumen of the artery is an area devoid of collagen fibrils and surrounded by multifenestrated sheaths that were previously the basal laminae of the hyaloid artery wall. Posteriorly, Cloquet’s canal opens into a funnel-shaped region anterior to the optic disc, known as the area of Martegiani.

Within the adult human vitreous fine, parallel fibers course in an anteroposterior direction, are continuous, and do not branch. The fibers arise from the vitreous base, where they insert anterior and posterior to the ora serrata. Various concepts are used to explain the connection between the peripheral anterior vitreous fibers and the retina and pars plana, but all agree that the pathophysiology of retinal tears is vitreous traction upon foci of strong adhesion at the vitreoretinal interface in these locations.

As the central fibers near the vitreous cortex course posteriorly, they are circumferential with the vitreous cortex, while central fibers “undulate” in a configuration parallel to Cloquet’s canal. Ultrastructural studies demonstrate that collagen, organized in bundles of packed, parallel fibrils, is the only microscopic structure that corresponds to these fibers. It is hypothesized that visible vitreous fibers form when hyaluronan molecules no longer separate the microscopic collagen fibrils, which results in the aggregation of collagen fibrils into bundles from which hyaluronan molecules are excluded. The areas adjacent to these large fibers have a low density of collagen fibrils and a relatively high concentration of hyaluronan molecules. Composed primarily of “liquid vitreous,” these areas scatter very little incident light and, when prominent, constitute “lacunae” seen in aging.

**Fig. 6.46-3** Vitreous anatomy according to classical anatomic and histological studies. (Reprinted with permission from Schepens CL, Neetens A, eds. The vitreous and vitreoretinal interface. New York: Springer-Verlag; 1987:20.)

**Fig. 6.46-4** The eye of a 57-year-old man after dissection of the sclera, choroid, and retina, with the vitreous still attached to the anterior segment. The specimen was illuminated with a slit-lamp beam shone from the side and the view here is at a 90° angle to this plane to maximize the Tyndall effect. The anterior segment is below and the posterior pole is at the top of the photograph. A large bundle of prominent fibers courses anteroposteriorly to exit via the premacular dehiscence in the vitreous cortex.

**Fig. 6.46-5** Human vitreous in old age. The central vitreous has thickened, tortuous fibers. The peripheral vitreous has regions devoid of any structure, which contain liquid vitreous. These regions correspond to “lacunae,” as seen clinically using biomicroscopy (arrows).
Microscopic Morphology

The vitreous cortex is defined as the peripheral “shell” of the vitreous that courses forward and inward from the anterior vitreous base, the “anterior vitreous cortex,” and posteriorly from the posterior border of the vitreous base, the “posterior vitreous cortex.” The posterior vitreous cortex is 100–110 µm thick and consists of densely packed collagen fibrils. Although no direct connections exist between the posterior vitreous and the retina, the posterior vitreous cortex is adherent to the internal limiting lamina of the retina, which is actually the basal lamina of retinal Müller cells. The exact nature of the adhesion between the posterior vitreous cortex and the internal limiting lamina is not known, but it most probably results from the action of various extracellular matrix molecules.

A hole in the prepapillary vitreous cortex can sometimes be visualized clinically when the posterior vitreous is detached from the retina (Fig. 6-46-6). If peripapillary glial tissue is torn away during posterior vitreous detachment and remains attached to the vitreous cortex about the prepapillary hole, it is referred to as Vogt’s or Weiss’ ring. Vitreous can extrude through the prepapillary hole in the vitreous cortex but does so to a lesser extent than through the premacular vitreous cortex. Various vitreomacularopathies can result. Other mechanisms, particularly tangential vitreomacular traction, are implicated in the pathogenesis of macular holes.

Embedded within the posterior vitreous cortex are hyalocytes. These mononuclear cells are spread widely apart in a single layer situated 20–50 µm from the internal limiting membrane of the retina. The highest density of hyalocytes is in the vitreous base, followed next by the posterior pole, with the lowest density at the equator. Hyalocytes are oval or spindle shaped, 10–15 µm in diameter, and contain a lobulated nucleus, a well-developed Golgi complex, smooth and rough endoplasmic reticula, many large lysosomal granules (periodic acid-Schiff positive), and phagosomes (Fig. 6-46-7). Balazs pointed out that hyalocytes are located in the region of highest hyaluronan concentration and suggested that these cells are responsible for hyaluronan synthesis. Hyalocyte capacity to synthesize collagen was first demonstrated by Newsome et al. Thus, in a similar fashion to the chondrocyte metabolism in the joint, hyalocytes may be responsible for vitreous collagen synthesis at some point(s) during life. The phagocytic capacity of hyalocytes is consistent with the presence of pinocytic vesicles and phagosomes and the presence of surface receptors that bind immunoglobulin G and complement. It is intriguing to consider that hyalocytes are among the first cells to be exposed to any migratory or mitogenic stimuli during various disease states, particularly proliferative vitreoretinopathy. Therefore, the role of these cells must be considered when the pathophysiology of all proliferative disorders at the vitreoretinal interface is considered, including premacular membrane formation.

The basal laminae about the vitreous are composed of type IV collagen closely associated with glycoproteins. At the pars plana, the basal lamina has a true lamina densa. The basal lamina posterior to the ora serrata is the internal limiting lamina of the retina. The layer immediately adjacent to the Müller cell is a lamina rara, which is 0.03–0.06 mm thick. The lamina densa is thinnest at the fovea (0.01–0.02 mm) and disc (0.07–0.1 mm). It is thicker elsewhere in the posterior pole (0.5–3.2 mm) than at the equator or vitreous base. The anterior surface of the internal limiting lamina (vitreous side) is normally smooth,
whereas the posterior aspect is irregular, as it fills the spaces created by the irregular surface of the subjacent retinal glial cells. This feature is most marked at the posterior pole, whereas in the periphery both the anterior and posterior aspects of the internal limiting lamina are smooth. The significance, if any, of this topographic variation is not known. At the rim of the optic disc the retinal internal limiting lamina is irregular, whereas the lamina of the optic nerve head, and is composed only of glycosaminoglycans with no collagen. This structural arrangement is known as the “central mesenchyma of Kuhn.” The thickness and chemical composition of these structures may account for, among other phenomena, the frequency with which abnormal cell proliferation arises from or near the optic disc in proliferative diabetic retinopathy and premacular membranes with macular pucker.

The vitreous is known to most firmly attach at the vitreous base, disc, and macula and over retinal blood vessels. The posterior aspect (retinal side) of the internal limiting lamina demonstrates irregular thickening the farther posteriorly from the ora serrata. So-called attachment plaques between the Müller cells and the internal limiting lamina have been described in the basal and equatorial regions of the fundus but not in the posterior pole, except for the fovea. It has been hypothesized that these develop in response to vitreous traction upon the retina. The thick internal limiting lamina in the posterior pole dampens the effects of this traction, except at the fovea, where the internal limiting lamina is thin. The thinness of the internal limiting lamina and the purported presence of attachment plaques at the central macula could explain the predisposition of this region to changes induced by traction. An unusual vitreoretinal interface overlies retinal blood vessels. Physiologically, this may provide a shock-absorbing function to dampen arterial pulsations. However, pathologically, this structural arrangement could also account for the proliferative and hemorrhagic events upon retinal blood vessels that are associated with vitreous traction.

**AGE-RELATED CHANGES**

**Embryology and Postnatal Development**

Early in embryogenesis, the vitreous is filled with blood vessels, the vasa hyaloidea propria. It is not known what stimulates regression of this hyaloid vascular system, but studies have identified a protein known to vitreous that inhibits angiogenesis in experimental models. Teleologically, this seems necessary not only to induce regression of the vascular primary vitreous but also to inhibit subsequent cell migration and proliferation and thereby minimize light scatter and achieve transparency. Identifying the phenomena inherent in this transformation may reveal how to control pathologic neovascularization in the eye and elsewhere.

**Developmental Anomalies**

Persistent fetal vascular (PFV) syndrome is an uncommon developmental anomaly in which the hyaloid vascular system of the primary vitreous fails to involute. This was initially described in detail by Reese in his 1955 Jackson Memorial Lecture, at which time the term employed was persistent hyperplastic primary vitreous (PHPV). Goldberg revisited this condition in his 1997 Jackson Memorial Lecture and coined the term PFV. There is a spectrum of PFV severity, ranging from pupillary strands and a Mittendorf’s dot to a dense retrolenticular membrane and/or retinal detachment. Anterior PFV consists of retro-lenticular fibrovascular tissue that attaches to the ciliary processes and lenticular fibrovascular tissue that attaches to the ora serrata. Anterior PFV consists of a prominent vitreous fibrovascular stalk that emanates from the optic nerve and the orbit of collagen and hyaluronan, whereas the posterior aspect is irregular, as it fills the spaces created by the irregular surface of the subjacent retinal glial cells. This feature is most marked at the posterior pole, whereas in the periphery both the anterior and posterior aspects of the internal limiting lamina are smooth. The significance, if any, of this topographic variation is not known. At the rim of the optic disc the retinal internal limiting lamina is irregular, whereas the lamina of the optic nerve head, and is composed only of glycosaminoglycans with no collagen. This structural arrangement is known as the “central mesenchyma of Kuhn.” The thickness and chemical composition of these structures may account for, among other phenomena, the frequency with which abnormal cell proliferation arises from or near the optic disc in proliferative diabetic retinopathy and premacular membranes with macular pucker.

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**Aging of the Vitreous**

Substantial rheological, biochemical, and structural alterations occur in the vitreous during aging. After 45–50 years of age a significant decrease occurs in the gel volume and an increase in the liquid volume of human vitreous. These findings were confirmed qualitatively in postmortem studies of dissected human vitreous, and liquefaction was observed to begin in the central vitreous. Vitreous liquefaction actually begins much earlier than the ages at which clinical examination or ultrasonography detects changes. Postmortem studies found evidence of liquid vitreous in eyes at 4 years of age and observed that by the time the human eye reaches its adult size (age 14–18 years) approximately 25% of the total vitreous volume consists of liquid vitreous. In these developing retinas, it was observed that after the age of 40 years a steady increase occurs in liquid vitreous simultaneously with a decrease in gel volume. By 80–90 years of age more than half the vitreous is liquid. The finding that the central vitreous is where fibers are first observed is consistent with the concept that breakdown of the normal hyaluronan-collagen association results in the simultaneous formation of liquid vitreous and aggregation of collagen fibrils into bundles of parallel fibrils, seen as large fibers (see Fig. 6-46-4). In the posterior vitreous such age-related changes often form large pockets of liquid vitreous, recognized clinically as lacunae, and mistakenly described as anatomic structures.

The mechanism of vitreous liquefaction is not understood. Gel vitreous can be liquefied in vivo through the removal of collagen by enzymatic destruction of the collagen network. Endogenous liquefaction may be the result of changes in the minor glycosaminoglycans and chondroitin sulfate profile of vitreous. It has been shown that the injection of chondroitinase ABC can induce liquefaction and “disinser-tion” of the vitreous. Plasmin is another agent being developed as an adjunct to vitreoretinal surgery because its ability to induce liquefaction of the central vitreous and desiccation at the vitreoretinal interface. Another possible mechanism of vitreous liquefaction is a change in the conformation of hyaluronan molecules with aggregation or cross-linking of collagen molecules. Singlet oxygen can induce conformational changes in the tertiary structure of hyaluronan molecules. Free radicals generated by metabolic and photosensitized reactions could alter hyaluronan and/or collagen structure and trigger a chain reaction that finally results in liquefaction. This is plausible because the cumulative effects of a lifetime of daily exposure to light may influence the structure and interaction of collagen and hyaluronan molecules by the proposed free radical mechanism(s).
confirmed posterior migration of the posterior border of the vitreous base during aging and also demonstrated intraretinal synthesis of collagen fibrils that penetrate the internal limiting of the retina and “splice” with vitreous collagen fibrils. These aging changes at the vitreous base could contribute to increased traction on the peripheral retina and to the development of retinal tears and detachment.

**Posterior Vitreous Detachment**

The most common age-related event in the vitreous is PVD. True PVD can be defined as a separation between the posterior vitreous cortex and the internal limiting lamina of the retina; PVD can be localized, partial, or total (up to the posterior border of the vitreous base). Autopsy studies reveal that the incidence of PVD is 63% by the eighth decade, and it is more common in myopic eyes, in which it occurs on average 10 years earlier than in emmetropic and hyperopic eyes. Cataract extraction in myopic patients introduces additional effects, which caused PVD to develop in all but 1 of 103 myopic (greater than −6 D) eyes.

Rheologic changes within the vitreous produce liquefaction, which, in conjunction with weakening of the vitreous cortex-internal limiting laminar adhesion, results in PVD. It is likely that dissolution of the posterior vitreous cortex–internal limiting laminar adhesion at the posterior pole allows liquid vitreous to enter the retrocortical space via the preapipillary hole and perhaps also the premacular vitreous cortex. With rotational eye movements, liquid vitreous can dissect a plane between the vitreous cortex and the internal limiting lamina, which results in true PVD. This volume displacement from the central vitreous to the preretinal space causes the observed collapse of the vitreous body (syneresis). Glare may be induced by PVD because of light scattering by the dense collagen fibril network in the posterior vitreous cortex.

“Floaters” are the most common complaint of patients with PVD. These usually result from entoptic phenomena caused by condensed vitreous fibers, glial tissue of epipapillary origin (which adheres to the posterior vitreous cortex), and/or intravitreal blood. Floaters move with vitreous displacement during eye movement and scatter incident light, which casts a shadow on the retina that is perceived as a gray, “hair-like” or “fly-like” structure. In 1935, Moore described “light flashes” as a common complaint that results from PVD. Wise noted that light flashes occurred in 50% of cases at the time of PVD; they were usually vertical and temporally located. Voerhof suggested that the light flashes result from the impact of the detached posterior vitreous cortex upon the retina during eye movement.

**Anomalous Posterior Vitreous Detachment**

Anomalous posterior vitreous detachment (APVD) occurs when the extent of vitreous liquefaction exceeds the degree of weakening of vitreoretinal adherence and traction is exerted at this interface. There are various causes for an imbalance between the degree of gel liquefaction and weakening of vitreoretinal adhesion. As described above, inborn errors of collagen metabolism, such as those present in Marfan’s, Ehlers-Danlos, and Stickler’s syndromes, result in extreme gel liquefaction at an early age when there is persistent vitreoretinal adherence. The result is a high incidence of large retinal tears and detachments. Systemic conditions such as diabetes induce biochemical and structural alterations in vitreous. These changes, referred to as diabetic vitreopathy, are important in the pathobiology of proliferative diabetic vitreoretinopathy and perhaps some cases of macular edema as well. Diabetic vitreopathy may one day be detected in vivo using noninvasive optical instrumentation. There are also changes associated with myopia, known as myopic vitreopathy, where there is excess vitreous liquefaction and a high incidence of large retinal tears and detachments.

Regardless of the underlying cause, abnormal traction at the vitreoretinal interface can have deleterious effects upon retina as well as vitreous (Table 6-46-1).

**Retinal effects**

Effects upon the retina vary, depending on the site affected. These include hemorrhage, retinal tears and detachment, and vitreomacular traction syndromes, including macular holes and some cases of diabetic macular edema. Proliferative diabetic retinopathy (PDR) can be aggravated by anomalous PVD. Lindner found that vitreous hemorrhage occurred in 13–19% of patients with PVD. Because vitreous hemorrhage results from considerable vitreoretinal traction, this finding in a patient with PVD is generally considered to be an important risk factor.
for the presence of a retinal tear and detachment. One 15-year study in Belgium found that in 126 cases of non-diabetic, nontraumatic vitreous hemorrhage that did not clear for 6 months, 25% were found to have retinal tears and 8% had retinal detachments. Another study found that in 36 eyes with fundus-obscuring vitreous hemorrhage, 24/36 eyes (67%) were found to have at least one retinal break, with 88% of breaks located in the superior retina. Eleven eyes (31%) had more than one retinal break. Fourteen of 36 eyes (39%) had a rhegmatogenous retinal detachment.

Retinal tears not involving blood vessels result from traction on other foci of usual vitreoretinal adhesion, such as the posterior border of the vitreous base. Abnormal foci of firm vitreoretinal adhesion, such as lattice degeneration and rosettes, are also frequently associated with retinal tears after PVD. Indeed, Byer has claimed that as many as 25% of the general population have some form of abnormal focal vitreoretinal adhesion, placing them at considerable risk from anomalous PVD.

**Vitreous effects**
Effects upon vitreous primarily involve posterior vitreoschisis. This condition results from splitting of the posterior vitreous cortex, with forward displacement of the anterior portion of the posterior vitreous cortex leaving part, or all, of the posterior layer of the split vitreous cortex still attached to the retina. Vitreoschisis has been detected in cases of proliferative diabetic vitreoretinopathy and retinovascular diseases, and likely has a role in the pathophysiology and sequelae of these conditions. Premacular membranes with macular pucker and cases of macular holes may also result from persistent attachment of partial or full-thickness posterior vitreous cortex to the macula while the remainder of the vitreous detaches forward. In the former case, tractional forces are centripetal (inward toward the fovea) causing macular pucker. In the latter condition tangential traction occurs in a centrifugal (outward from the fovea) direction, causing a macular hole.

**METABOLIC DISORDERS OF VITREOUS**

**Diabetic Vitreopathy**
In humans who have diabetes, there is an increase in vitreous glucose levels. These elevated levels of glucose are associated with increased nonenzymatic glycation products in human vitreous collagen and elevated levels of the enzyme-mediated cross-link dihydroxylysylornitireine. Also, considerable diabetic effects may involve hyaluronan. In the daily management of diabetes, significant fluctuations in the systemic concentrations of a variety of molecules may occur, which can alter the ionic milieu of the vitreous. Shifts in systemic metabolism, and in turn osmolarity and hydration of the vitreous, could result in periodic swelling and contraction of the entire vitreous, with consequent traction upon structures attached to the posterior vitreous cortex, such as new blood vessels that have grown out of the optic disc and/or retina. These events could influence the course of diabetic retinopathy as they may contribute to the proliferation of neovascular frond and perhaps even induce rupture of the new vessels and cause vitreous hemorrhage.

**Asteroid Hyalosis**
This generally benign condition is characterized by small yellow-white spherical opacities throughout the vitreous. The prevalence of asteroid hyalosis was previously found to be 0.042–0.5%, although a recent study of 10,801 autopsy eyes found an incidence of 1.96%; it affects all races but with a male-to-female ratio of 2:1. Curiously, asteroid hyalosis is unilateral in over 75% of cases. Asteroid bodies are associated intimately with the vitreous gel and move with typical vitreous displacement during eye movement, which suggests a relationship with collagen fibril degeneration. However, PVD, either complete or partial, occurs less frequently in individuals with asteroid hyalosis than in age-matched controls, which does not support age-related degeneration as a cause. Histological studies demonstrate a crystalline appearance and a pattern of positive staining to fat and acid mucopolysaccharide stains.

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**TABLE 6-46-1 ANOMALOUS POSTERIOR VITREOUS DETACHMENT**

<table>
<thead>
<tr>
<th>Traction Site(s)</th>
<th>Retinal Effects</th>
<th>Vitreous Effects</th>
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<tbody>
<tr>
<td>Blood vessels</td>
<td>Retinal hemorrhages</td>
<td>Vitreous hemorrhage</td>
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<tr>
<td></td>
<td>Aggravate retinal neovascularization</td>
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<tr>
<td>Macula</td>
<td>Vitreomacular traction syndrome</td>
<td>Vitreoschisis with macular pucker</td>
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<tr>
<td></td>
<td>Diabetic macular edema (diffuse)</td>
<td>Macular holes</td>
</tr>
<tr>
<td>Periphery</td>
<td>Retinal tears/detachments</td>
<td>White without pressure</td>
</tr>
<tr>
<td>Optic disc</td>
<td>Vitreopapillary traction syndrome</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Aggravate NVD (PDVR, CRVO)</td>
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</tbody>
</table>

CRVO, central retinal vein occlusion; NVD, neovascularization of the optic disc; PDVR, proliferative diabetic vitreoretinopathy.
that is not affected by hyaluronidase pretreatment. Electron diffraction studies showed the presence of calcium oxalate monohydrate and calcium hydroxypophosphate. Ultrastructural studies reveal intertwined ribbons of multilaminar membranes (with a 6 nm periodicity) that are calcium hydroxyphosphate. Ultrastructural studies reveal intertwined calcium phosphate crystals.

Some reports suggest an association between asteroid hyalosis and diabetes mellitus, while other investigations found no such association. Asteroid hyaloid appears to be the main element in asteroid bodies. Electron diffraction structural analysis demonstrated calcium hydroxyapatite and possibly other forms of calcium phosphate crystals.

Amyloidosis

Amyloidosis can result in the deposition of opacities in the vitreous of one or both eyes. Bilateral involvement can be an early manifestation of the dominant form of familial amyloidosis, although rare cases of vitreous involvement in multifocal forms have been reported. The opacities first appear in the vitreous adjacent to retinal blood vessels and later appear in the anterior vitreous. Initially, the opacities are granular with wispy fringes and later take on a “glass wool” appearance. When the opacities form strands, they appear to attach to the retina and the posterior aspect of the lens by thick footplates. Following PVD, the posterior vitreous cortex is observed to have thick, linear opacities that follow the course of the retinal vessels. The opacities seem to aggregate by “seeding” on vitreous fibrils and along the posterior vitreous cortex.

Histopathological specimens contain star-like structures with dense, fibrillar centers. The amyloid fibrils are 5–10 nm in diameter and are detectable from the vitreous body to the vitreous surface, and by the fact that the vitreous fibrils are very straight and long. Electron microscopy can confirm the presence of amyloid, and immunocytochemical studies identified the major amyloid constituent as a protein that resembles praealbumin. Streeten proposed that hyalocytes could perform the role of macrophage processing of the amyloid protein before its polymerization. This may further explain why the opacities initially appear at the posterior vitreous cortex where hyalocytes reside.

REFERENCES
