Invisible (Fig. 3.1) by design, vitreous was long unseen as important in the physiology and pathology of the eye. Recent studies have determined that vitreous plays a significant role in ocular health (1) and disease (1,2), including a number of important vitreoretinal disorders that arise from abnormal embryogenesis and development. Vitreous embryology is presented in detail in Chapter 1. Notable is that primary vitreous is filled with blood vessels during the first trimester (Fig. 3.2). During the second trimester, these vessels begin to disappear as the secondary vitreous is formed, ultimately resulting in an exquisitely clear gel (Fig. 3.1). The following will review vitreous development and the congenital disorders that arise from abnormalities in hyaloid vessel formation and regression during the primary vitreous stage and biochemical abnormalities related to secondary vitreous dysgenesis.

**BIOCHEMISTRY OF THE VITREOUS BODY**

At birth, vitreous is composed primarily of collagen resulting in a dense appearance on dark-field slit microscopy (Fig. 3.3), because collagen scatters light intensely. Hyaluronan (HA) synthesis begins after birth. Due to considerable hydrophilicity, HA generates a “swelling” pressure within the burgeoning vitreous that contributes to growth of the eye and also spreads apart the collagen fibrils to minimize light scattering inducing transparency (Figs. 3.1 and 3.4). This is evident when comparing dark-field microscopy of a human embryo (Fig. 3.3) with similar imaging in a 4-year-old child (Fig. 3.4) (see also Chapter 4).

There is heterogeneous distribution of collagen throughout the vitreous. Chemical (3,4) and light-scattering studies (5) have shown that the highest density of collagen fibrils is present in the vitreous base, followed by the posterior vitreous cortex anterior to the retina, and then the anterior vitreous cortex behind the posterior chamber and lens. The lowest density is found in the central vitreous and adjacent to the anterior cortical gel. HA molecules have a different distribution from collagen, being most abundant in the posterior cortical gel with a gradient of decreasing concentration centrally and anteriorly (6,7).

Both collagen and HA are synthesized during childhood. Total collagen content in the vitreous gel remains at about 0.05 mg until the third decade (8). As collagen does not appreciably increase during this time but the size of the vitreous does increase with growth, the density of collagen fibrils effectively decreases. This could potentially weaken the collagen network and destabilize the gel. However, since there is active synthesis of HA during this time, the dramatic increase in HA concentration may stabilize the thinning collagen network (9).

Bishop (10) proposed that the leucine-rich repeat protein, opticin, is the predominant structural protein in short-range spacing of collagen fibrils. Scott (11) and Mayne et al. (12) claimed that HA plays a pivotal role in stabilizing the vitreous gel via long-range spacing. However, studies (13) using HA lyase to digest vitreous HA found that the gel structure was not destroyed, suggesting that HA is not essential for the maintenance of vitreous gel stability, leading to the proposal that collagen alone is responsible for the gel state of vitreous (10).
Section I | Development of the Eye and Retina

Transforming growth factor (TGF) has also been implicated in ocular vasculature remodeling (23). Since all four forms of TGF-β were found in vitreous hyalocytes, these cells may be involved in the induction of hyaloid vessel regression. Other mechanisms may be involved. There is evidence that family members of the Wnt signaling pathway, particularly Wnt7b expressed in macrophages, are involved in normal hyaloid regression (24) in the mouse, although this mechanism has not been confirmed in humans.

SECONDARY VITREOUS

Secondary (avascular) vitreous formation arises from a remodeling of the earlier primary (vascular) vitreous (25). By the 40- to 60-mm stage in humans, the VHP has evidence of regression is apparent as early as 11 WG (16). The hyaloid system shows clear signs of regression by 13 to 15 WG beginning in the VHP followed by the TVL and then the PM (16–18).

Although the precise biochemical process involved in hyaloid vasculature formation and regression is poorly defined, studies have shown that vascular endothelial growth factor (VEGF) may trigger the growth of the TVL and PM (19–22). Thus, decreased VEGF levels may induce vascular regression. Transforming growth factor (TGF) has also been implicated in ocular vasculature remodeling (23). Since all four forms of TGF-β were found in vitreous hyalocytes, these cells may be involved in the induction of hyaloid vessel regression. Other mechanisms may be involved. There is evidence that family members of the Wnt signaling pathway, particularly Wnt7b expressed in macrophages, are involved in normal hyaloid regression (24) in the mouse, although this mechanism has not been confirmed in humans.

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reached maximum maturity and regression begins (22). The posterior VHP is the first component to degenerate. The regression continues to move centrally affecting the TVL and finally the hyaloid artery itself. Although flow through the hyaloid artery is decreasing during this time, ceasing entirely around the 240-mm stage, the artery may continue to elongate as the eye lengthens (26). As the primary vitreous regresses, its remnants provide a framework for collagen fibrils of the secondary vitreous that form via interactive remodeling. As this occurs, the primary and secondary vitreous coexist in the same space during this transitional developmental period (25,26).

**VITREOUS STRUCTURE**

In the human embryo, vitreous structure (27–33) has a dense homogenous appearance (Fig. 3.3) primarily as a result of the aforementioned collagen synthesis during secondary vitreous formation. HA synthesis after birth separates collagen fibrils, inducing transparency (Fig. 3.4). In the absence of diabetes and myopia, vitreous remains largely transparent until around the fifth decade when fine, parallel fibers appear coursing in an anteroposterior direction (Fig. 3.5B and C) (2,30–32). The fibers arise from the vitreous base (Fig. 3.5H) where they insert anterior and posterior to the ora serrata. As the peripheral fibers course posteriorly, they are circumferential with the vitreous cortex, while central fibers “undulate” in a configuration parallel with Cloquet canal (33). The fibers are continuous and do not branch. Posteriorly, these fibers insert into the vitreous cortex (Fig. 3.5E and F).

Ultrastructural studies (34) have demonstrated that collagen fibrils are the only microscopic structures that could correspond to these fibers. These studies also detected the presence of bundles of packed, parallel collagen fibrils. Eventually, the aggregates of collagen fibrils attain sufficiently large proportions so as to be visualized in vitro (Fig. 3.5) and clinically. The areas adjacent to these large fibers have a low density of collagen fibrils separated by HA molecules and therefore do not scatter light as intensely as the bundles of collagen fibrils.

**Vitreous Base**

Vitreous base is a three-dimensional zone extending 1.5 to 2 mm anterior to the ora serrata, 1 to 3 mm posterior to the ora serrata (35), and several millimeters into the vitreous body itself (36). The posterior extent of the posterior border of the vitreous base varies with age (37,38). In a variety of vitreoretinal developmental disorders, the temporal vitreous never forms; thus, there is a different peripheral terminus bounded by vitreous gel posteriorly and liquid vitreous anteriorly (see below). Vitreous fibers enter the vitreous base by splaying out to insert anterior and posterior to the ora serrata (Fig. 3.5H). The anterior-most fibers form the “anterior loop” of the vitreous base, a structure that is important in the pathophysiology of anterior proliferative vitreo-retinopathy (2,39–41). Studies by Gloor and Daicker (42) showed that cords of vitreous collagen insert into gaps between the neuroglia of the peripheral retina. They likened this structure to Velcro (a self-adhesive nylon material) and proposed that this would explain the strong vitreoretinal adhesion at this site. In the anterior vitreous base, fibrils interdigitate with a reticular complex of fibrillar basement membrane material between the crevices of the nonpigmented ciliary epithelium (43). The vitreous base also contains intact cells that are fibroblast-like anterior to the ora serrata and macrophage-like posteriorly. Damaged cells in different stages
of involution and fragments of basal laminae, presumed to be remnants of the embryonic hyaloid vascular system, are also present in the vitreous base (43).

**Vitreous Cortex**

The vitreous cortex is the peripheral shell of the vitreous body that courses forward and inward from the anterior vitreous base to form the anterior vitreous cortex and posteriorly from the posterior border of the vitreous base to form the posterior vitreous cortex. The anterior vitreous cortex, often referred to as the “anterior hyaloid face,” begins about 1.5 mm anterior to the ora serrata. The posterior vitreous cortex is 100 to 110 µm thick (44) and consists of densely packed collagen fibrils (Fig. 3.6, bottom). There is no vitreous cortex over the optic disc (Fig. 3.4A), and the cortex is thin over the macula due to rarefaction of the collagen fibrils (44). The prepapillary hole in the vitreous cortex can sometimes be visualized clinically when the posterior vitreous is detached from the retina. If peripapillary glial tissue is torn away during posterior vitreous detachment (PVD) and remains attached to the vitreous cortex around the prepapillary hole, it is referred to as Vogt or Weiss ring. Vitreous can extrude through the prepapillary hole in the vitreous cortex (Fig. 3.4A) but does so to a much lesser extent than through the premacular vitreous cortex where it can produce traction and certain forms of maculopathy (45–47).

**Hyalocytes**

Hyalocytes are mononuclear cells (Fig. 3.7) that are embedded in the vitreous cortex and widely spread apart in a monolayer 20 to 50 µm from the retina. Quantitative studies of cell density in the bovine (48) and rabbit (49) vitreous found the highest density of hyalocytes in the vitreous base, followed by the posterior pole, and the lowest density at the equator. Hyalocytes are oval or spindle-shaped, are 10 to 15 µm in diameter, and contain a lobulated nucleus, a well-developed Golgi complex, smooth and rough endoplasmic reticula, and many large periodic acid-Schiff–positive lysosomal granules and phagosomes (44,50). Balazs (51) pointed out that hyalocytes are located in the region of highest HA concentration and suggested that these cells are responsible for vitreous HA synthesis. There is evidence to suggest that hyalocytes maintain ongoing synthesis and metabolism of glycoproteins within the vitreous (52,53). Hyalocytes have also been shown to synthesize vitreous collagen (54) and enzymes (55).

The phagocytic capacity of hyalocytes has been described in vivo (56) and demonstrated in vitro (57,58), consistent with the presence of pinocytic vesicles and phagosomes (50) and surface receptors that bind immunoglobulin G and complement (58). Hyalocytes become phagocytic cells in response to inducting stimuli and inflammation. HA may have a regulatory effect on
Vitreoretinal Interface

In the child, it is virtually impossible to mechanically detach the posterior cortical vitreous from the retina, resulting in surgical complications such as retinal breaks. The interface between vitreous and retina consists of a complex formed by the posterior vitreous cortex and the internal limiting lamina (ILL), which includes the basal lamina of Müller cells (62,63) (Fig. 3.6) (see also Chapter 4). The ILL is composed of type IV collagen closely associated with hyalocytes (64–66). Immediately adjacent to the Müller cells is the lamina rara, which is 0.03 to 0.06 µm thick and demonstrates no species variations nor changes with topography or age. The lamina densa is thinnest at the fovea (0.01 to 0.02 µm). It is thicker elsewhere in the posterior pole (0.5 to 3.2 µm) than at the equator or vitreous base. At the rim of the optic nerve head, the retinal ILL ceases, although the basement membrane continues as the inner limiting membrane of Elschnig (67). This membrane is 50 nm thick and is believed to be the basal lamina of the astroglia in the optic nerve head. At the central-most portion of the optic disc, the membrane thins to 20 nm, follows the irregularities of the underlying cells of the optic nerve head, and is composed only of glycosaminoglycans and no collagen (68). This structure is known as the central meniscus of Kuhnt. Balazs (7) has stated that the Müller cell’s basal lamina prevents the passage of cells as well as molecules larger than 15 to 20 nm. Consequently, the thinness and chemical composition of the central meniscus of Kuhnt and the membrane of Elschnig may account for, among other effects, the frequency with which abnormal cell proliferation arises from or near the optic nerve head (39,41).

Zimmerman and Straatsma (69) claimed that there are fine, fibrillar attachments between the posterior vitreous cortex and the ILL and proposed that this was the source for the strong adhesion between vitreous and retina. The composition of these fibrillar structures is not known, and their presence has never been confirmed. It is more likely that an extracellular matrix “glue” exists between the vitreous and retina in a fascial as opposed to focal apposition (30,64,65), comprised of fibronectin, laminin, and other extracellular matrix components (70). Western blots of separated proteins derived from dissected vitreoretinal tissues identified two chondroitin sulfate–containing proteoglycans of approximately 240-kDa molecular weight (MW) that are believed to function as adhesive molecules at the vitreoretinal interface. These findings formed the rationale for experimental and clinical studies on pharmacologic vitreolysis (71) using ABC chondroitinase (see Chapter 2.2).

Development of the Embryonic Hyaloid Vasculature

Many developmental vitreoretinal disorders result from abnormalities in the embryonic vascular system of the vitreous (VHP) and lens (TVL). These attain maximum prominence during the 9th week of gestation or 40-mm stage (72). Hyaloid vessel development around the seventh WG appears to be driven by VEGF165 (an isoform of VEGF-A). However, once the hyaloid vascular system reaches completion, the alternatively spliced antiangiogenic VEGF165b isoform begins to dominate (73).

Regression of Fetal Vasculature

Atrophy of the vessels begins posteriorly with dropout of the VHP, followed by the TVL. Recent studies have detected the onset of apoptosis in the endothelial cells of the TVL as early as day 17.5 in the mouse embryo (19). At the 240-mm stage (7th month) in the human, blood flow in the hyaloid artery ceases. Regression of the vessel itself begins with glycogen and lipid deposition in the endothelial cells and pericytes of the hyaloid vessels (74). Endothelial cell processes then fill the lumen, and macrophages form a plug that occludes the vessel. The cells in the vessel wall then undergo necrosis and are phagocytized by mononuclear phagocytes (75) identified as hyalocytes (76).

Recent studies suggest that the VHP and the TVL regress via apoptosis (77). Mitchell et al. (19) point out that the first event in hyaloid vessel regression is endothelial cell apoptosis and propose that lens development separates the fetal vasculature from VEGF-producing cells, decreasing the levels of this survival factor for vascular endothelium, inducing apoptosis. Following endothelial cell apoptosis, there is loss of capillary integrity, leakage of erythrocytes into the vitreous, and phagocytosis of apoptotic endothelium by hyalocytes. Meeson et al. (78) proposed that there are actually two forms of apoptosis that are important in regression of the fetal vitreous vasculature. The first (“initiating apoptosis”) results from macrophage induction of apoptosis in a single endothelial cell of an otherwise healthy capillary segment with normal blood flow. The isolated dying endothelial cells project into the capillary lumen and interfere with blood flow. This stimulates synchronous apoptosis of downstream endothelial cells.
remain unclear and more research in this area is needed. Often lack comparison or controls, so mechanisms that EIF2 signaling and protein ubiquitination were the top canonical pathways (82). The presence of dystroglycan and profilin-1 was confirmed in the hyaloid vessels of 10, 14, and 18 WG embryonic human eyes by immuno-labeling (80). Clusterin was not present in the HA or TVL of 10 and 14 WG eyes, but was found in the HA of 18 WG eyes. Cadherin was not found in the hyaloid vessels of the 10 WG eyes but was observed in the hyaloid vessels at 14 and 18 WG (80).

Comparison of the protein composition of the adult human and developing human vitreous from the second trimester (14 to 20 WG) revealed 314 and 1,213 proteins, respectively, including 78 proteins that were unique to the adult and 1,002 unique to the fetal vitreous (81). A large fraction of the fetal vitreous proteins are intracellular and may be involved in the remodeling of the hyaloid vasculature and retina during the second trimester. Proteins that were found in greater abundance in fetal vitreous compared to adult vitreous include low MW kininogen-1, cystatin-B, insulin-like growth factor–binding protein (IGFBP)-2 and IGFBP-4, thrombospondin-1 and thrombospondin-4, thyredoxin, mu-crystallin, alpha-2-antiplasmin, nidogen-2 and growth differentiation factor 8, as well as secreted proteins pigment epithelial derived factor (PEDF), AMBP (α-1-microglobulin), hemoglobin, and Ig chains. Low MW extracellular matrix proteins detected in fetal vitreous included COL5A1, 6A3, 11A1, 15A1, 1A1, 2A1, 4A2, and 18A1 (81).

Bioinformatic analysis of the protein profile of fetal (14 to 20 WG) human vitreous revealed that most proteins were members of either the free radical scavenging, molecular transport, or small molecule biochemistry, or connective tissue disorders networks. Those proteins involved in free radical scavenging, molecular transport, and small molecule biochemistry were all intracellular and decreased during the second trimester. In contrast, the proteins in the connective tissue disorders network were all extracellular and increased during the second trimester. This pattern is consistent with replacement of the cellular primary vitreous with the acellular collageneous secondary vitreous. This analysis further revealed that EIF2 signaling and protein ubiquitination were the top canonical pathways (82).

These studies, essential because they are in humans, often lack comparison or controls, so mechanisms remain unclear and more research in this area is needed.

Pathologies of the Primary Vitreous
Persistence of the hyaloid vascular system occurs in 3% of full-term infants but in 95% of premature infants (83). 90% of infants born earlier than 36 WG, and in over 95% of infants weighing <5 pounds at birth (84). There is a spectrum of disorders resulting from persistence of the fetal vasculature (85), which can at times be associated with preapillary hemorrhage (86).

a. Mittendorf dot is a remnant of the anterior fetal vascular system located at the former site of anastomosis of the hyaloid artery and TVL. It is usually inferonasal to the posterior pole of the lens and is not associated with any known dysfunction.

b. Bergmeister papilla is the occluded remnant of the posterior portion of the hyaloid artery with associated glial tissue. It appears as a gray, linear structure anterior to the optic disc and adjacent retina and does not cause any known functional disorders. Exaggerated forms can present as preapillary veils.

c. Vitreous cysts are generally benign lesions that are found in eyes with abnormal regression of the anterior (87) or posterior (88) hyaloid vascular system, otherwise normal eyes (89,90), and eyes with coexisting ocular disease, such as retinitis pigmentosa (91) and uveitis (92). Some vitreous cysts contain remnants of the hyaloid vascular system (93), supporting the concept that the cysts result from abnormal regression of these embryonic vessels (94). However, one histologic analysis of aspirated material from a cyst purportedly revealed cells from the retinal pigment epithelium (95). Vitreous cysts are generally not symptomatic and thus do not require surgical intervention. However, there is a report (96) describing the use of Nd:YAG laser therapy to rupture a free-floating posterior vitreous cyst.

d. Persistent fetal vasculature (PFV) (persistent hyperplastic primary vitreous, PHPV) was first described by Reese (97) and subsequently renamed PFV (85). Though there is a broad spectrum of presentations, PHPV/PFV often features a plaque of retrolental fibrovascular connective tissue adherent and posterior to the posterior lens capsule. The membrane extends laterally to attach to the ciliary processes, which are elongated and displaced centrally. Although 90% of cases are unilateral, many of the fellow eyes had Mittendorf dot or another anomaly of anterior vitreous development (98). A persistent hyaloid artery, often still perfused with blood, arises from the posterior aspect of the retrolental plaque in the affected eye. In severe forms, there can be microphthalmos with anterior displacement of the lens–iris diaphragm, shallowing of the anterior chamber, and secondary glaucoma. PHPV/PFV is believed to arise from abnormal regression and hyperplasia of the primary
vitreous (97). Experimental data suggest that the abnormality begins at the 17-mm stage of embryonic development (99). The hyperplastic features result from proliferation of retinal astrocytes and a separate component of glial hyperplasia arising from the optic nerve head (100). The fibrous component of the PHPV/PFV membrane is presumably synthesized by these cells (101). A recent case report found that collagen fibrils in this fibrous tissue had diameters of 40 to 50 nm with a cross-striation periodicity of 65 nm. The investigators concluded that the collagen fibrils differed from those of the primary vitreous and suggested that they arose either from a different population of cells or were the result of abnormal metabolism by the same cells that synthesize vitreous collagen (102).

The retina is usually not involved in anterior PHPV, suggesting that the anterior form is due to a primary defect in lens development and that vitreous changes are all secondary (103). In posterior PHPV/PFV opaque connective tissue arises from Bergmeister papilla and persistent hyaloid vessels (101,104). These can cause congenital falciform folds of the retina and, if severe, can cause tent-like retinal folds, leading on rare occasions to tractional and/or rhegmatogenous retinal detachment. Font and investigators (105) demonstrated the presence of adipose tissue, smooth muscle, and cartilage within the retrolental plaque and suggested that PHPV/PFV arises from metaplasia of mesenchymal elements in the primary vitreous.

e. Familial (Dominant) Exudative Vitreoretinopathy

Familial exudative vitreoretinopathy (FEVR) was first described in 1969 by Criswick and Schepens (106) as a bilateral, slowly progressive abnormality that resembles retinopathy of prematurity (ROP), but without a history of prematurity or postnatal oxygen administration (see also Chapter 28 on FEVR). Gow and Oliver (107) identified this disorder as an autosomal dominant condition with complete penetrance. They characterized the course of this disease in stages ranging from PVD with snowflake opacities (stage I) to thickened vitreous membranes and elevated fibrovascular scars (stage II) and vitreous fibrosis with subretinal and intraretinal exudates, ultimately developing retinal detachment due to fibrovascular proliferation arising from neovascularization in the temporal periphery (stage III). Plager et al. (108) recently reported the same findings in four generations of three families but found X-linked inheritance. van Nouhuys (109–111) studied 101 affected members in 16 Dutch pedigrees and five patients with sporadic manifestations. He found that the incidence of retinal detachment was 21%, with all but one case occurring prior to the age of 30. These were all tractional or combined traction–rhegmatogenous detachments, and there were no cases of exudative retinal detachment. van Nouhuys (109) concluded that the etiology of FEVR lies in premature arrest of development in the retinal vasculature, since the earliest findings in these patients were nonperfusion of the peripheral temporal retina with stretched retinal blood vessels and shunting with vascular leakage. Thus, van Nouhuys considers FEVR as a retinopathy with secondary vitreous involvement. However, Brockhurst et al. (112) reported that vitreous membrane formation began just posterior to the ora serrata and that it preceded retinal vessel abnormalities, suggesting a vitreous origin to this disorder. Others suggested that there may be a combined etiology involving anomalies of the hyaloid vascular system, primary vitreous, and retino-vascular dysgenesis (113).

f. Retinopathy of Prematurity:

First described in the 1940s (114), retinopathy of prematurity remains a leading cause of blindness in children in the United States (115). Vitreous liquefaction, often unidentified (116) in stages 1 and 2 ROP, likely occurs as a result of reactive oxygen species (117) but also overlying the peripheral retina where immature Müller cells may not support typical gel synthesis and may account for the vitreous trough apparent during surgery for stage 4 ROP. The disrupted composition may limit the inherent vitreous ability to inhibit cell invasion (118–120), thereby permitting neovascularization in stage 3 ROP to grow (121,122) between posterior gel and peripheral liquid vitreous anteriorly (Fig. 3.8) (122). Instability at the interface between gel and liquid vitreous exerts...
traction on the underlying ridge. As ROP advances from stage 3 to 4, the neovascular tissue grows through the vitreous along the walls of the future Cloquet canal or the tractus hyaloideus of Eisner toward Wiegert ligament on the posterior lens capsule (123). Primary vitreous cells, responding to intraocular angiogenic stimuli, likely proliferate and migrate to help create the dense central vitreous stalk and retrolental membrane seen in the cicatricial stages.

Pathologies of the Secondary Vitreous

Vitreous Collagen Disorders: As vitreous is one of many connective tissues in the body, it is of interest to consider parallel phenomena occurring in the vitreous and connective tissues elsewhere, especially as related to collagen. Long ago, Gärtner (124) pointed out the similarities between the intervertebral disk and the vitreous, in which age-related changes with herniation of the nucleus pulposus were associated with presenile vitreous degeneration in 40% of cases. He proposed that a generalized connective tissue disorder resulted in disk herniation and presenile vitreous degeneration in these cases. Based on these findings, Gärtner likened herniation of the nucleus pulposus in the disc to prolapse of vitreous into the retrohyaloid space by way of the posterior vitreous cortex following PVD (Fig. 3.4D).

Maumenee (125) identified several different disorders with single-gene autosomal dominant inheritance in which dysplastic connective tissue primarily involves joint cartilage. In these conditions, there is associated vitreous liquefaction, collagen condensation, and vitreous syneresis (collapse). Since type II collagen is common to cartilage and vitreous, Maumenee suggested that various arthroophthalmopathies might result from different mutations, perhaps of the same or neighboring genes, on the chromosome involved with type II collagen metabolism. In these disorders, probably including such conditions as Wagner disease (126), the fundamental problem in the posterior segment of the eye is that the vitreous is liquefied and unstable and becomes synergetic at an early age. Vitreous syneresis (collapse) occurs in the vitreoretinal interface in concert with the changes inside the vitreous body. Thus, in these cases, abnormal type II collagen metabolism causes destabilization of the vitreous and results in traction on the retina that can lead to large posterior tears and difficult retinal detachments.

i. **Marfan Syndrome.** In Marfan syndrome, an autosomal dominant disorder featuring poor musculature, lax joints, aortic aneurysms, and arachnodactyly, there is lens subluxation, thin sclera, peripheral fundus pigmentation changes, and vitreous liquefaction at an early age. Myopia, vitreous syneresis, and abnormal vitreoretinal adhesions at the equator likely account for the frequency of rhegmatogenous retinal detachment caused by equatorial or posterior horseshoe tears (129). Marfan syndrome is associated with mutations in the FBN1 gene that encodes the connective tissue protein fibrillin 1.

ii. **Ehlers-Danlos Syndrome**

*Systemic Manifestations:* Ehlers-Danlos syndrome has some similarities to Marfan syndrome, most notably joint laxity, aortic aneurysms, and an autosomal dominant pattern of inheritance. However, there are as many as six types of Ehlers-Danlos patients, and both autosomal dominant and autosomal recessive forms have been described. A further distinction from Marfan patients is hyperelastic skin and poor wound healing of all connective tissues, including cornea and sclera. Ehlers-Danlos syndrome has been associated with mutations in numerous collagen genes, including COL1A1, COL1A2, COL3A1, COL5A1, and COL5A2.

*Ocular Manifestations:* Ocular manifestations include lens subluxation, angioid streaks, thin sclera, and high myopia due to posterior staphyloma. Vitreous liquefaction and syneresis occur at a young age. Vitreous traction causes vitreous hemorrhage, perhaps also due to blood vessel wall fragility, and retinal tears with rolled edges, often causing bilateral retinal detachments (129).

iii. **Stickler Syndrome**

*Genetics:* In 1965, Stickler et al. (130) described a condition in five generations of a family that was found to be autosomal dominant with complete penetrance and variable expressivity. Stickler syndrome is the most common type II/XI collagenopathy, arising from mutations in at least three collagen genes. Although typically inherited in an autosomal dominant fashion, families with an autosomal recessive pattern of inheritance have been described (131). Stickler syndrome is most commonly associated with mutation in COL2A1, a 54-exon–containing gene coding for type II collagen. In one instance (132,133) of a family with typical Stickler syndrome, posterior chorioretinal atrophy and vitreoretinal degeneration were found, even though they have been classically associated with Wagner disease. Vitreous findings from that study validated reports that mutations in the COL2A1 gene result in an optically empty vitreous with retrolenticular membrane phenotype.

*Systemic Manifestations:* In general, the features of Stickler syndrome were a marfanoid skeletal habitus and orofacial and ocular abnormalities. Subsequent studies identified subgroups with short stature and a Weill-Marchesani habitus. The skeletal abnormalities now accepted as characteristic of Stickler syndrome are radiographic evidence of flat epiphyses, broad metaphyses, and especially
spondyloepiphyseal dysplasia (134). In general, the systemic manifestations can vary greatly, even within a family all possessing the same genotype. Distinguishing features can include deafness, cleft palate, joint hypermobility, and premature arthritis (135). Recently, however, an ocular-only subgroup has been identified, which has no systemic features but a high risk of rhegmatogenous retinal detachments (136,137).

Ocular Manifestations: Ocular abnormalities are high myopia, >−10 Da in 72% of cases (138), and vitreoretinal changes characterized by vitreous liquefaction, fibrillar collagen condensation, quadrandric lamellar cortical lens opacities (135), and a perivascular lattice-like degeneration in the peripheral retina believed to be the cause of a high incidence (>50%) of retinal detachment (134). There is some evidence of phenotype/genotype correlation, which has led to the classification of Stickler syndrome patients into five subgroups (135). Additionally, the vitreous can exhibit three distinct phenotypes, membranous (type 1), beaded (type 2), or normal (139). Patients with abnormalities in the genes coding for type II procollagen and type XI α1 procollagen are the ones who have severe vitreous abnormalities. Patients with type XI α2 procollagen defects typically present without ocular manifestations (140). Another study (141) analyzed the ultrastructural feature of a vitreous membrane with multiple fenestrations in a patient with a Stickler syndrome. A type 2 vitreous phenotype was found in the left eye, whereas the other eye’s vitreous abnormalities appeared to result from a conversion to a type 1 phenotype. In such a conversion, a fenestrated membrane may represent the posterior vitreous cortex in a complete PVD. The fenestrated membrane is made of avascular fibrocellular tissue with cells arranged cohesively around the fenestration. Results from ultrastructural findings were characteristic of proliferating Müller cells, and collagen fibrils were shown to be similar to normal vitreous by ultrastructural examination. The authors concluded that collagen molecules are not functionally modified, but they are probably quantitatively insufficient during vitreous development.

iv. Knobloch Syndrome: Knobloch (142) described an autosomal recessive syndrome similar to Stickler syndrome with hypotonia, relative muscular hypoplasia, and mild to moderate spondyloepiphyseal dysplasia causing hyperextensible joints. The vitreoretinopathy is characterized by vitreous liquefaction, veils of vitreous collagen condensation, and perivascular lattice-like changes in the peripheral retina. Retinal detachment in patients with Knobloch syndrome has been explained by loss-of-function mutations in COL18A1, the gene encoding the α1 chain of collagen XVIII based on findings from one investigation (143) demonstrating that collagen XVIII is crucial for anchoring vitreous collagen fibrils to the inner limiting membrane.

v. Myopia: It has been proposed (144) that myopia unrelated to the aforementioned arthroophthalmopathies should also be considered a disorder of vitreous collagen. The anomalous PVD (127,128) that results from extensive liquefaction of vitreous (myopic vitreopathy) and propensity for retinal detachment due to peripheral retinal traction and myopic peripheral retinal degeneration suggest that this postulate deserves closer scrutiny. Indeed, vitreomacular traction has already been identified as an important component in the pathogenesis of myopia-related pathologies, including myopic foveal retinoschisis (145).

Although the exact mechanism for increased vitreous liquefaction in high myopia is unclear, several theories have been proposed. Dysfunctional Müller cell activity is an older hypothesis that derived from studies showing abnormal B wave results on electroretinograms of highly myopic patients (Current thinking is that the B-wave does not arise from Mueller cells but perhaps from ON-center bipolar cells) (146). However, the absence of abnormally thick inner limiting laminae in myopic patients (147) suggests that Müller cells may not be the culprit. Another study showed that highly myopic patients with ocular pathology (macular detachments or macular hole) had increased vitreous and serum levels of transthyretin (TTR) compared to controls (148). Additionally the TTR in the macular detachment cases was abnormally stable suggesting a misfolded protein. TTR is a homotetrameric protein that functions as a carrier for both thyroxin and retinol-binding protein and has previously been implicated in several amyloid-related diseases such as vitreous amyloidosis, Alzheimer disease, and familial amyloidotic polyneuropathy (148,149). These results suggest that TTR may be a biomarker of myopic vitreopathy while also playing a role in the pathophysiology of myopic ocular conditions.

REFERENCES


