Episcleral Venous Pressure and Glaucoma

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BACKGROUND

The pioneering work of Lauber in the early 20th century was the first stepping stone towards understanding the role of episcleral venous pressure in aqueous humor dynamics. This led to Seidel’s observation in 1923 that India ink injected in the anterior chamber gained access to episcleral veins, and in 1942 Ascher made the first direct observation of the aqueous veins.

EPISCLERAL VENOUS PRESSURE AND AQUEOUS OUTFLOW

Aqueous outflow via the conventional outflow pathway is dependent on the pressure gradient between intraocular pressure (IOP) and episcleral venous pressure (EVP). According to the Goldmann equation: IOP = F/C + EVP (F aqueous humor inflow, C outflow facility). This implies that EVP elevation uncompensated by reduced aqueous inflow or increased aqueous outflow facility will result in increased IOP. The definite relation between EVP and IOP, however, is unclear. The normal episcleral venous pressure is 8-10 mm Hg; variability depends on the measurement technique. It is thought that for every mm Hg rise in EVP, there is an equal increase in the intraocular pressure; although it has been suggested that the magnitude of rise of intraocular pressure can be more.

EFFECT OF ANGIOARCHITECTURE ON EVP

The episcleral vasculature shows morphological specialities that distinguish it from other vessels. There are only few capillaries, but numerous arteriovenous anastomoses and an extended network of venules with a wall containing circularly arranged muscle cells. Furthermore there is a strikingly dense vasoconstrictive and vasodilative innervation of the veins. Animal studies show that the elaborated innervation of the episcleral anastomoses is a prerequisite for a subtle modulation of the blood flow and possibly of the aqueous humor outflow dynamics. However, the role of the episcleral vasculature and its specialities for regulation of IOP as well as possible therapeutic aspects are still not fully understood and require further investigation.

EFFECT OF POSTURE ON EVP AND IOP

EVP has been shown to be influenced by body posture. In a study, mouse EVP was successfully measured based on the detection of erythrocyte reflux from an episcleral vein into Schlemm’s canal. Both EVP and IOP increased with the degree of the head-down body position. EVP and IOP remain relatively constant throughout the day. It increases in the supine position with a corresponding increase in IOP. Another study has shown no difference in the intraocular pressure and episcleral venous pressure in the younger and older group in the sitting position, but they were higher in the supine position for older subjects.

EVP IN NORMAL AND GLAUCOMATOUS EYES

EVP in normal eyes and in eyes with open angle glaucoma have not be shown to be significantly different. Kupfer reported that ocular hypertensive eyes have a slightly lower EVP than normal eyes, but his group was small for a statistical comparison. Talusan also found that EVP of the patients with ocular hypertension were found to be significantly lower than those of the patients with glaucoma and the normal patients. Other investigators noted that the EVP/intraocular pressure (IOP) ratio was significantly different in NTG patients (80.0% +/-3.2) in comparison with both POAG patients (67.1% +/- 2.8) and controls (69.2% +/- 2.4). Regression analysis revealed a significant linear correlation between EVP and IOP in both the NTG and the POAG group. In the control group, however, the correlation was weak.

CAUSES OF ELEVATED EVP

Causes of elevated EVP may be divided into three different groups: venous obstruction, which includes thyroid ophthalmopathy, retrobulbar tumor, cavernous sinus or orbital vein thrombosis, episcleral or orbital vein vasculitis, and...
obstruction of the superior vena cava; arteriovenous anomalies, including carotid artery—cavernous sinus fistula, orbital varices, dural shunts and Sturg-Weber syndrome; and idiopathic. Ocular laterality, motility dysfunction and fundoscopic signs of venous congestion are dependent upon the level of blockage.

**CLINICAL PRESENTATION**

**Idiopathic Elevated Episceral Venous Pressure (IEEVP)**

The idiopathic type of elevation of EVP is a diagnosis of exclusion after intraorbital and intracranial pathology has been excluded. The diagnosis is based on the clinical findings of prominent (arterialized) episcleral veins (Fig. 1), elevated IOP causing characteristic optic nerve and/or visual field changes typical of glaucoma and an open angle on gonioscopy. Diagnostic evaluation should include a complete ophthalmic examination, radiological imaging such as an MRI and non-invasive vascular imaging to exclude a cerebrovascular disorder. Idiopathic episcleral venous stasis with secondary open angle glaucoma (Radius Maumenee syndrome) responds to microsurgical sinusotomy to reduce the pressure gradient from Schlemmm’s canal to episcleral vessels.

**Venous Obstruction**

Elevated EVP may be present in severe cases of thyroid ophthalmopathy with marked proptosis and orbital congestion (Fig. 2). Lesions of upper thorax may obstruct venous return from the head and produce exophthalmos, edema and cyanosis of the face.

**Arteriovenous Fistulas**

Carotidocavernous sinus fistulas can be either spontaneous or traumatic. The mixing of arterial and venous pressure leads to a reduction in arterial pressure and an elevation in EVP. The pattern of venous drainage, either anterior into the ophthalmic veins or posterior into the petrosal sinuses, often dictates the clinical findings and radiographic appearance. Anterior drainage typically leads to the most dramatic ocular findings and enlargement of the superior orbital vein, the latter often detectable with CT or MRI. These patients have prominent episcleral and conjunctival vessels along with elevated IOP and closure of the fistula is the primary condition required for favorable IOP control. Arteriovenous anomalies may be treated primarily with intravascular embolization and balloon occlusion. Newer techniques using transvenous approaches have also shown promising outcome.

**Orbital Varices**

Orbital varices typically have intermittent episodes of raised EVP associated with valsalva maneuver. Since EVP is normal between episodes, glaucoma is not common.

**Sturge-Weber Syndrome**

IOP elevation may occur due to the episcleral hemangiomas with arteriovenous fistulas. Phelps proposed that the veins draining aqueous from the canal of Schlemm are a part of an intrascleral or episcleral hemangioma or the canal of Schlemm itself may be part of the hemangioma. Arteriovenous shunts in the hemangioma raise episcleral venous pressure, which in turn elevates intraocular pressure. Blood may be seen in Schlemm’s canal on gonioscopy but it is not pathognomonic (Fig. 3). In glaucoma associated with Sturge-Weber syndrome, medical treatment often fails to control intraocular pressure, thus requiring surgical intervention that may result in serious complications.

Goniotomy, trabeculotomy, combined trabeculotomy-trabeculectomy, drainage devices and nonpenetrating filtering surgery have been tried with variable results.
Choroidal effusion and postoperative serous retinal detachment are possible complications. Moderately tight scleral flap suture, posterior sclerostomy at the time of filtering surgery and delayed suture lysis may be required to prevent choroidal effusion. Drainage implants cannot be used in cases with severe proptosis and orbital congestion due to space limitations.

**Orbital Amyloidosis**

Rare cases of localized orbital amyloidosis with secondary glaucoma due to elevated EVP have been reported. Nelson et al postulate that perivascular infiltration of amyloid around extraocular vessels may cause elevated EVP.

**Diagnosis and Management**

Theses include ultrasound biomicroscopy, orbital ultrasonography, computerized axial tomography, magnetic resonance imaging and angiography. The finding of a hyperintense signal in the cavernous sinus on MRA source imaging provides additional, and sometimes the only, neuroradiographic Carotico-cavernous fistula evidence. In addition a team approach involving a glaucomatologist, neuro-ophthalmologist, neuroradiologist and an orbital surgeon are often necessary for accurate diagnosis and appropriate management.

Treatement of underlying cause of elevated EVP is ideal, but may not be always possible. Miotics and laser trabeculoplasty are not very helpful due to normal facility of outflow in these patients. Aqueous suppressants may help in some patients. Surgical management holds the key for controlling elevated intraocular pressure, but sudden decompression should be avoided as these are prone to choroidal effusion. However, even if the underlying pathology is identified and treated, the IOP likely may remain elevated if the venous system has had sufficient time to arteriolize.

**REFERENCES**


“...The wonderful beauty of prayer is that the opening of our hearts is as natural as the opening of a flower. To let a flower open and bloom, it is only necessary to let it be, so if we simply are, if we become and remain still and silent, our heart cannot but be open, the spirit cannot but pour through into our whole being.”

—John Main