Retinal Venous Occlusive Disease

Retinal vein occlusion is an eye condition commonly seen by retinal specialists. It is second only to diabetic retinopathy as a cause of visual loss due to retinal vascular disease. There are two forms of retinal vein occlusion, branch retinal vein occlusion (BRVO) and central retinal vein occlusion (CRVO). While there are similarities in the pathogenesis and clinical nature of these two events, each has unique etiologies, differential diagnosis, management and prognosis.

Branch Retinal Vein Occlusion

A branch retinal vein occlusion is essentially a blockage of the portion of the circulation that drains the retina of blood. The arteries deliver blood to the retina. The red blood cells and plasma then course through the capillaries and eventually into the venous system, beginning with small veins and ending with larger ones, and eventually reaching the central retinal vein. With blockage of any vein, there is back–up pressure in the capillaries, which leads to hemorrhages and also to leakage of fluid and other constituents of blood. Usually, the occlusion occurs at a site where an artery and vein cross. The occlusion site determines the extent or distribution of the hemorrhage, ranging from a small vein branch to a quadrantic occlusion involving one fourth of the retina to a hemispheric (hemi–retinal) occlusion involving one half of the retina to an occlusion of the central retinal vein, which involves the entire retina (when the central vein is involved, this is called a central retinal vein occlusion which is discussed below).

Branch retinal vein occlusions are by far the most common cause of retinal vascular occlusive disease. Males and females are affected equally. Most occlusions occur after age 50, although younger patients are sometimes seen with this disorder (in this age group it is often called papillophlebitis). The highest rate of occurrence is in individuals in their 60’s and 70’s. The risk factors for this disorder are similar to those for vascular occlusive disease elsewhere in the body such as stroke and coronary artery disease. Specifically, aging, high blood pressure, diabetes, and smoking are all risk factors. Glaucoma has also been identified as a risk factor in some studies. There are less common conditions which may put a patient at risk for developing a vein occlusion including blood clotting abnormalities such as hyperhomocysteinemia, activated protein C resistance (Factor V Leiden), protein C and S deficiency, anti–phospholipid antibodies and diseases which cause sludging of the circulation or so–called hyperviscosity. Inflammatory and infectious conditions which cause vasculitis such as sarcoidosis and tuberculosis are also risk factors for vein occlusion. In general, unless there is a reason to suspect these less common conditions (such as young age, history of previous thrombophilia, or history suggestive of inflammation or infection), exhaustive laboratory testing is usually not indicated. Most patients are referred to their internist for the appropriate medical evaluations.

The diagnosis of a retinal branch vein occlusion poses little difficulty to an ophthalmologist who will detect dilated blood vessels, hemorrhages, and swelling (edema) in the distribution of the vein. It appears that the more complete the blockage,
the more intense the hemorrhages and the edema. In fact, the blockage may be so
dramatic that the involved capillaries cease to function and close off (ischemia or
capillary non–perfusion). About 10% of patients suffering from a branch vein occlusion
will experience a branch or a central vein occlusion in the fellow eye in the future.

There are three complications of branch retinal vein occlusion which threaten vision:
macular edema, macular ischemia (non–perfusion) and neovascularization (growth of
new abnormal blood vessels).

When the distribution of the vein involves the center of the retina (macula), bleeding and
exudation or leakage occurs there, producing symptoms. Leakage in the macula causes
macular edema in which a patient will have blurred vision and loss of portions of the
field of vision (corresponding to the distribution of the obstructed vein). Basically, the
edema damages the architecture of the retina, causing these symptoms. These visual
changes can be monitored with an Amsler grid. A fluorescein angiogram and OCT may
be useful in evaluating macular edema and determining whether treatments with laser
or pharmacological therapies are necessary.

In the first three to six months after the occurrence of the branch vein occlusion, there is
often significant intraretinal hemorrhage that involves the macula, making it difficult to
predict the clinical course and visual outcome. After the first few months, it may be
useful to do a fluorescein angiogram and OCT. Fluorescein angiography is helpful in
analyzing the retinal circulation, particularly the capillaries which may manifest
abnormalities such as leakage or macular ischemia (non–perfusion: closure of blood
vessels which supply the retina with oxygen and other nutrients). OCT is useful
for detecting retinal swelling (edema).

If the fluorescein angiogram indicates that capillary non–perfusion is the cause of the
vision loss, it is unlikely that the vision will improve significantly over time. However, if
the poor vision is due to edema or swelling, laser photocoagulation or pharmacological
therapies, such as steroids or anti–vasogenic drugs (eg. Avastin), may be useful in
sealing leaking capillaries to enhance resolution of the edema for stabilization and
improvement of the vision. Intraocular steroids, which are commonly used as a
treatment for macular edema, increase the chance of cataract formation and elevation
of intraocular pressure (steroid–induced glaucoma).

Sometimes in venous occlusive disease, scar tissue can form on the surface of the
retina. This condition, which is called a macular pucker or an epiretinal membrane may
result in distorted vision (metamorphopsia) which is not improved with laser or
pharmacologic treatment. Vitrectomy surgery may be indicated for the removal of a
macular pucker.

The most devastating potential problem in a vein occlusion is that of neovascularization.
The neovascularization may develop in 40% of those cases where branch vein
occlusions produce large areas of capillary non–perfusion. This retinal
neovascularization generally develops in the first 6 to 12 months after the occlusion.
Unless laser treatment is performed, at least 60% of the patients with neovascularization will experience episodes of vitreous hemorrhage. In severe cases of neovascularization, retinal detachment can occur from pulling by these vessels and associated scar tissue on the retina (traction detachment).

Laser photocoagulation treatment is a proven therapy for neovascularization in vein occlusions. Indeed, laser treatment can cause stabilization or, at times, regression of the vascular growth. This treatment, while important in helping to prevent further visual loss, is not usually associated with improvement in vision. As vein occlusions evolve, some normal vessels may dilate to compensate for the obstructed vein. Sometimes, these collateral vessels may be difficult to distinguish from neovascularization on clinical examination. A fluorescein angiogram may be useful in this determination.

Recently, the intraocular injection of anti-vasogenic drugs, that tell blood vessels to stop growing, has shown promising results in the control of retinal neovascularization. These treatments have not yet been assessed in prospective clinical trials but may be used on an off-label basis. One particular medication that has been increasingly utilized for this purpose is Avastin, a drug that is approved for the intravenous treatment of colon cancer. Intraocular injections of Avastin have shown promising early results, and an excellent safety profile, in the control of retinal swelling and neovascularization due to a variety of retinal conditions. Avastin lasts about 6 weeks in the eye after a single injection and may need to be repeated if the disease reactivates. Although an injection into the eye sounds painful, it is simple to perform, relatively painless, and very well tolerated by patients.

There is no known medical treatment for retinal branch vein occlusion. Anti–coagulants such as heparin, coumadin and aspirin have not been shown to be of value in preventing branch vein occlusion or managing its complications. Because anti–coagulants may be associated with systemic complications, they are prescribed only in specific clinical circumstances, for example for patients with known clotting abnormalities.
Central Retinal Vein Occlusion

Central retinal vein occlusion is closure of the final retinal vein (located at the optic nerve) which collects all of the blood after it passes through the capillaries. The systemic risk factors for branch retinal vein occlusion mentioned above are also risk factors for central retinal vein occlusion.

Central retinal vein occlusion is generally categorized into two forms: non–ischemic and ischemic. This means that some central retinal vein occlusions are associated with a significant obstruction of capillaries or non–perfusion. This predisposes to a peculiar type of neovascularization that occurs in front of the eye on the iris (rubeosis irides). These eyes may develop a very high pressure known as neovascular glaucoma due to obstruction of the fluid outflow channels. This is a very serious complication which is associated with severe vision loss and may cause pain and loss of the eye itself. Laser photocoagulation treatment is very useful in managing rubeosis irides. If performed early in the course (when iris neovascularization is first detected), it may help prevent these complications. Patients with recent central retinal vein occlusions must be followed frequently in order to detect this complication in a timely manner.

Less frequently than in branch vein occlusion, patients with central retinal vein occlusion, may also develop neovascularization in the back of the eye, causing vitreous hemorrhage and retinal detachment. Laser treatment may be useful in managing these complications.

As with branch retinal vein occlusion, macular edema and non–perfusion are also frequently seen with central retinal vein occlusion. Macular edema, even without significant macular ischemia, is not treated routinely with laser photocoagulation. This is because a recent study failed to show a benefit for patients with central retinal vein occlusion, particularly for those who are elderly. (In contrast, laser treatment has been shown to be effective for patients with branch retinal vein occlusion). It is possible, but not proven, that some young patients with central vein occlusion of the non–ischemic type may benefit from localized laser treatment for macular edema.

Some patients with macular edema from central vein occlusions may respond to the off–label intraocular injection of steroids (triamcinolone) or anti–vasogenic drugs (Avastin). These pharmacologic agents wear off over a period of approximately two months and may need to be re–injected if the edema returns. Optical Coherence Tomography (OCT) is a useful test for the detection of macular edema in CRVO and for following the response to treatment.

If the fluorescein angiogram indicates that capillary non–perfusion is the cause of the vision loss, it is unlikely that the vision will improve significantly over time regardless of any treatment.
If a patient develops an occlusion of the central vein in both eyes, there is a greater possibility of an underlying systemic cause. It is recommended (by our group) that such patients have a thorough medical work-up as outlined previously.

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In summary, retinal vein occlusions develop from obstruction of the venous outflow from the eye. The blockage may vary in size and location, accounting for a wide range of retinal outcomes. Some of the complications of retinal vein occlusion may be appropriately managed with laser treatment. It is hoped that through further research, even better strategies for prevention and management will be developed.