Management of ocular ischaemic syndrome

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Symptoms of carotid artery disease frequently present to ophthalmologists. Though these may be sight threatening, they may represent the first signs of life threatening carotid artery stenosis. These include cerebral transient ischaemic attacks (TIA), transient monocular blindness (amaurosis fugax), central or branch retinal artery occlusion, hypotensive retinopathy (previously known as "venous stasis retinopathy"), and ocular ischaemic syndrome (OIS). Of these associations with carotid artery disease, OIS presents the most challenging condition for the ophthalmologist with many controversial aspects to its management.

OIS is a severe form of chronic ischaemia of both anterior and posterior segments of the eye as well other orbital structures supplied by the ophthalmic artery. It is thought to be due to chronic hypoperfusion when carotid artery stenosis is greater than 90%.

OIS is rare; however, ophthalmoscopic features of hypertensive retinopathy have been found in 5%-20% of patients with carotid artery occlusive disease and approximately 200 patients with OIS have been reported in the literature by way of case reports, retrospective reviews, and prospective studies. Based on a questionnaire survey it has been estimated that a neuro-ophthalmologist or glaucoma specialist encounters at least one case per year.

In this article we present an overview and recommendations for best clinical practice for OIS based on an extensive review of recent studies.

Clinical findings in OIS

SYMPTOMS
Clinical presentation may include sudden (41%), gradual (28%), or transient vision loss (15%) or pain, either ocular or orbital (13%). In 20% of cases, the clinical signs of OIS are an incidental, asymptomatic finding. Brown et al. in their retrospective study of 43 patients, showed that 90% had reduced visual acuity and in two thirds of patients this occurred gradually over a period of weeks to months.

Rarely, vision loss may be precipitated by exposure to bright lights ("bright light amaurosis") with subjective after-images of visual distortion, fragmentation, dazzle, or just blurring. It may also occur with change in posture or even exertion. It is likely that this phenomenon represents photoreceptor ischaemia due to poor retinal and choroidal circulation. Russell and Ikeda showed that patients with carotid artery stenosis whose main symptom was unilateral bright light amaurosis had marked ERG delay in the recovery time of b-wave amplitude after photostress. These patients also had associated fluorescein fundus angiogram (FFA) changes of patchy choroidal filling.

Ocular discomfort or pain around the orbit in OIS, in the absence of glaucoma, occurs in 5-10% of patients. It often reduces on lying down and is thought to be due to ischaemic damage to the branches of the ophthalmic division of the trigeminal nerve.

SIGNS
Anterior segment signs include dilated episcleral vessels, corneal oedema, anterior chamber cells, and pronounced flare ("ischaemic pseudo-inflammatory uveitis"), a mid-dilated poorly reactive pupil, cataract, iris atrophy, iris neovascularisation with or without angle neovascularisation, or neovascular glaucoma.

Iris neovascularisation has been found in up to 90% of cases, and dilated episcleral vessels and uveitis in up to 20% of cases. Iris neovascularisation seen at presentation is considered an indicator of poor visual prognosis. Over 95% of such eyes develop a visual acuity of counting fingers or less within 1 year. Intraocular pressure (IOP) is usually raised. However, it may be normal or even reduced, despite iris neovascularisation, presumably due to ciliary body ischaemia leading to reduced aqueous production. Two thirds of patients will have an IOP of <22 mm Hg at presentation.

Posterior segment signs include venous dilatation with or without tortuosity, mid-peripheral retinal haemorrhages and microaneurysms, and an easily induced retinal artery pulsation with gentle digital pressure. Ischaemic changes include retinal arteriolar narrowing, retinal capillary non-perfusion, macular oedema, optic disc neovascularisation (NVD) and, less commonly, retinal neovascularisation (NVE). Easily inducible or even spontaneous retinal artery pulsation is present in most cases. It is a striking feature and most pronounced near the optic disc.

Severe carotid artery stenosis has also been suggested to explain asymmetry in proliferative diabetic retinopathy. In diabetic patients with OIS, in whom the anterior segment signs are mild or even absent, it may appear, therefore, as asymmetrical proliferative diabetic retinopathy.

Anterior ischaemic optic neuropathy has been reported in association with OIS as a rare complication of carotid artery obstruction. It is believed that this is due to an inadequate perfusion pressure within the deep capillaries of the optic nerve head.

DIFFERENTIAL DIAGNOSIS
It is important to exclude other important causes of iris neovascularisation such as proliferative diabetic retin-
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opathy and ischaemic central retinal vein occlusion (CRVO). Retinal arterial pressure should be normal in eyes with diabetic retinopathy and venous occlusion.

INVESTIGATIONS

Ocular

Fluorescein fundus angiography—The fluorescein angiographic signs of OIS include delayed and patchy choroidal filling, increased retinal arteriovenous circulation times, areas of retinal capillary non-perfusion, late leakage from arterioles and veins, leakage from new vessels, and macular oedema. Leakage from retinal vessels has been reported to be present in 85% of eyes with OIS, presumably due to ischaemic endothelial cell dysfunction. In combination with leakage from microaneurysms this appears to account for macular oedema when present.

With this in mind, during angiographic evaluation of macular oedema in routine practice, any delay in filling time should raise the possibility of coincidental carotid artery disease.

Retinal capillary non-perfusion is sometimes present in OIS. Brown and Magargal reviewed the FFA of 40 eyes of 40 patients and found retinal capillary non-perfusion in some cases, most often in the mid-periphery with a gradual transition. The authors did not specify any further details as to the frequency or extent of this finding. This contrasts with the sharply demarcated border between areas of retinal capillary perfusion and non-perfusion that are seen with, for example, an ischaemic central vein occlusion. However, Mizioń et al failed to show any retinal capillary non-perfusion, even diabetics, in 22 eyes from 15 patients suitable for FFA. The authors emphasise the point that in the absence of capillary non-perfusion the results of diabetic trials for proliferative diabetic retinopathy and therefore the beneficial value of pan-retinal laser photocoagulation (PRP) may not be applicable to OIS.

The role of FFA in OIS is therefore to aid confirmation of diagnosis, determine the cause of iris neovascularisation, and to demonstrate retinal capillary non-perfusion in order to validate the indication for PRP.

Orbital ophthalmic colour Doppler ultrasound—Colour Doppler imaging (CDI) can be useful. This is a recent advance in ultrasonography that provides colour coded blood flow data of vessels at the same time as conventional real time grey scale B-scan images. Retrolubar colour Doppler ultrasound findings in OIS with more than 70% carotid artery stenosis include reduced peak systolic velocities in ophthalmic and central retinal arteries and continuous or intermittent reversal of ophthalmic artery blood flow. However, colour Doppler imaging has limited clinical use at present because measurements of flow velocity in orbital vessels are poorly reproduced, the most reliable being those of the ophthalmic artery nasal to the globe.

General

Systemic diseases most often associated with OIS include diabetes mellitus (56%), hypertension (50–73%), ischaemic heart disease (38–48%), and cerebrovascular disease (27–31%). The prevalence of hypertension and ischaemic heart disease was comparable with those found in patients with retinal embolic disease. However, the prevalence of diabetes mellitus in OIS was not only greater than the aged matched population but also greater than in patients with retinal embolic disease. It is suggested that perhaps the combination of both large vessel and small vessel disease increases the risk of OIS.

Importantly, the 5 year mortality rate in OIS patients is as high as 40%. The majority of deaths are due to cardiac disease.

Management of OIS

Controversy in the management of OIS arises from the fact that the majority of patients reported in the literature are part of small retrospective series or case reports. Only one prospective study with 39 eyes of 32 patients has been published and the largest retrospective series collected only 52 cases over 8 years in an outpatient clinic that recorded 1.5 million visits during that period. With such a rare disease, it is very difficult to carry out a randomised prospective trial evaluating the effects of treatment on visual outcome.

ROLE OF THE OPHTHALMOLOGIST

Ocular treatment is directed towards the treatment of anterior segment inflammation, ablation of retinal ischaemia (if present), and the control of raised intraocular pressure and neovascular glaucoma (NVG).

It is important to diagnose OIS early. Intraocular pressure needs to be controlled and attention directed towards the prevention of NVG. To this purpose hypertensive retinopathy should be assessed and monitored both clinically and with FFA in order to identify retinal ischaemia and validate the need for effective laser PRP. In the acute stage it is important to arrange FFA as soon as possible before significant corneal oedema precludes a view for funduscopy and effective laser PRP.

Anterior segment inflammation may be treated with regular topical steroid and cycloplegics. Topical β adrenergic antagonists or α adrenergic agonists along with oral carbonic anhydrase inhibitors are first line therapy for raised IOP; however, they may only reduce IOP temporarily. Topical cyclopiaegia along with oral analgesics are required for pain relief.

Optic nerve function monitoring, including visual fields, is important as a guide to the progression of disease, the effect of treatment, and the presence of coincidental, treatable ocular disease.

It should not be forgotten that patients with bilateral disease will often be eligible for registration as blind or partially sighted by way of field loss before central vision is dramatically affected. Patients with reduced visual acuity will require low vision aids.

A prompt referral for full medical and neurological assessment in order to optimise systemic risk factors and associated cardiovascular disease is needed. Medical treatment would include aspirin or another antiplatelet drug, treatment of hypertension and diabetes, and advice to stop smoking. The decision regarding treatment for carotid artery disease requires both the neurologist and vascular surgeon.

ROLE OF PANRETINAL PHOTOCOAGULATION

Panretinal photocoagulation (PRP) is the accepted treatment for retinal ischaemia predisposing to neovascularisation in diabetic eye disease. It is thought that retinal ischaemia triggers the production of retinal angiogenic growth factors that stimulate retinal (NVE) and optic nerve head (NVD) new vessel growth and possibly diffuse into the anterior segment giving rise to iris neovascularisation (NVI). By ablating ischaemic retina, it is thought that PRP reduces the production of growth factors thereby leading to regression of neovascularisation thus preventing NVI.
In OIS, the occurrence of NVI has traditionally been attributed to severe retinal ischaemia. It has been found, however, that PRP alone may cause NVI regression in only 36% of eyes. Laser PRP is thought not to be as effective in reducing the ischaemic stimulus for NVI as for diabetic neovascularisation. In this context it is notable that Mizener et al. found no evidence of retinal ischaemia, in the form of capillary dropout on angiography in OIS patients, even in those with coincident diabetes mellitus. In animal studies Hayreh and Baines experimentally induced NVI due to uveal ischaemia in rhesus monkeys without any retinal ischaemia. It has been suggested therefore, that uveal ischaemia alone may be responsible for neovascularisation in some cases of OIS.

These observations highlight the importance of angiography in the investigation of OIS. It has been suggested that PRP in OIS should be reserved for cases of established retinal ischaemia. In this context full peripheral retinal ablation 3000–5000 burns of 200–500 µm spot size be used. Hayreh comments that there is no scientific rationale for PRP when FFA shows no retinal ischaemia in the form of capillary non-perfusion. In such cases of choroidal or ciliary body ischaemia rather than retinal ischaemia, the complications of PRP such as pain and further visual field constriction are therefore avoided.

CONTROL OF INTRAOCULAR PRESSURE

NVG is notoriously difficult to treat in cases of OIS. Chronic IOP elevation in the presence of compromised ocular perfusion can lead to anterior ischaemic optic neuropathy, central retinal artery occlusion, corneal oedema, and a blind painful eye.

Topical β-adrenergic antagonists or α adrenergic agonists, topical steroids, and cycloplegics along with oral carbonic anhydrase inhibitors may temporarily help in reducing IOP and inflammation. However, medical therapy is usually not effective in controlling IOP in the intermediate to long term because the trabecular meshwork is physically occluded with neovascular tissue and fibrosis. Attention should therefore also be directed towards eliminating any ischaemic retina present. Conventional filtering surgery (trabeculectomy usually with mitomycin C) also carries a limited chance of success in the presence of NVI. Rarely, tube shunt procedures (such as the Molteno tube or Ahmed valve implant) may be considered as a primary procedure or after failed conventional filtering surgery in a sighted eye.

Ciliary body ablation, however, has been shown to be effective in controlling IOP in end stage refractory glaucomas such as NVG. Methods such as cyclocryotherapy and laser cyclophotocoagulation are well described. Until recently the 1064 nm contact or non-contact neodymium-YAG (Nd:YAG) laser has been a popular modality. More recently, however, the 810 nm semiconductor diode laser has been shown to be better absorbed by ciliary body pigment and offers more effective cycloablation.

Diode laser (cyclodiode) ciliary ablation is reported to cause less inflammation than Nd:YAG laser, is less painful, and results in a more predictable final IOP. Complications such as phthisis bulbii, hypotony, and uveitis are also considered to be less frequent. A typical protocol strategy would be to apply approximately 10 laser burns of 1500–2000 mW for 1500–2000 ms in each quadrant. To titrate treatment with response, these laser applications may be applied over a number of treatment sessions. An appropriate end point would be long term control of intraocular pressure with clearing of corneal oedema. There may or may not be concomitant improvement in visual function. Clinical regression of NVI using peripheral trans-scleral retinal diode combined with cyclodiode is also reported.

ROLE OF CAROTID SURGERY

OIS is usually an important indicator of carotid artery stenosis, and all OIS patients should be referred for neurological and cardiovascular assessment at the time of ocular diagnosis. Carotid endarterectomy has been shown to benefit patients with symptomatic cerebral ischaemia when there is greater than 70% carotid artery stenosis.

Based on retrobulbar colour Doppler ultrasound examinations, carotid endarterectomy has been shown to improve ocular blood flow. Peak systolic velocity of flow in the ophthalmic artery rises after surgery and any reversal of ophthalmic artery flow is corrected. Carotid artery surgery therefore can reduce ocular ischaemia and improve hypotensive retinopathy as well as reduce the risk of stroke.

Although reports exist of IOP rising as ciliary body circulation is improved by carotid endarterectomy or by superficial temporal artery-middle cerebral artery bypass surgery (STA-MCA), most patients undergoing carotid endarterectomy do not experience any significant rise in IOP.

Clinical signs of hypotensive retinopathy have been reported to regress following carotid surgery. FFA changes following surgery include reduction in arteriovenous transit time, macular oedema, and microaneurysms present.

Surgery has also been shown in selected cases to help regression of iris neovascularisation and neovascular glaucoma.

It should be borne in mind that many of the cases reported above were of patients undergoing STA-MCA bypass surgery rather than carotid endarterectomy. Bypass surgery has been advocated if lesions are unresectable by carotid endarterectomy. Such a situation would include cases of total internal carotid stenosis, common carotid occlusion, or diffuse ulcerative stenosis extending distally along the internal carotid artery. More recently, an international randomised clinical trial of STA-MCA bypass surgery in patients with symptomatic carotid occlusive disease failed to show any protection against cerebral ischaemia with no reduction in stroke rate. STA-MCA bypass surgery is no longer a widely accepted alternative to carotid endarterectomy and, to some extent, the case for carotid artery surgery in OIS is still not proved. The European Carotid Surgery Trial (ECST) results showed that the risk of ischaemic stroke over 3 years in symptomatic patients with 70–99% carotid stenosis on medical treatment alone was only about 20%. They also showed that carotid endarterectomy lowered this risk by 50% over 3 years. Therefore, surgery had no benefit and possibly harmful effects in the remaining 80% of patients. A recent prognostic risk model based on the ECST proposed a “risk factor score” and suggested that patients with severe carotid stenosis and a recent cerebral rather than ocular event had a greater risk of stroke when taking medical treatment and therefore a greater benefit from surgery.

It should also be remembered that carotid surgery may not alter long term visual outcome in the affected eye.

Cases of early improved visual acuity after surgery have been reported and a small retrospective series reported stabilisation of visual acuity; however the authors did not report long term visual outcome, for example, at 1 year.

Conclusion

OIS is a severe but rare condition, often leading to significant visual loss and chronic ocular pain. Iris neovascularisation is an indicator of poor visual prognosis. Diagnosis
should aim to detect early disease before the onset of iris neovascularisation. Current medical and laser treatment protocols can successfully control progression of disease although treatment for vision is usually poor and it is difficult to make clear recommendations for aggressive therapy. It is important to recognise that the carotid circulation is significantly compromised in most cases of OIS and if life threatening neurological or cardiovascular complications are to be avoided, prompt neurological and medical advice should be sought.

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