Birdshot retinochoroidopathy and subretinal new vessels

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SUMMARY Decrease of visual acuity in birdshot retinochoroidopathy is due either to optic atrophy or to 3 types of macular involvement: cystoid macular oedema, geographic atrophy, or macular serous detachment. We describe 3 cases of juxtapapillary subretinal neovascularisation occurring in long-standing birdshot retinochoroidopathy. The mechanism of the formation of the new vessels is discussed.

Birdshot retinochoroidopathy was first described by Ryan and Maumenee in 1980.1 This bilateral syndrome is a chronic disease with exacerbations and remissions. It is characterised by bilateral vitreous inflammation with no ‘snow-bank’ exudates overlying the pars plana, diffuse retinal vasculitis, and multifocal patches of depigmentation of the choroid and of the pigment epithelium in the postequatorial fundi. These characteristic lesions led Gass2 to propose the term ‘vitiliginous chorioretinitis’ because of the association in some patients of this disorder with vitiligo. Papillary involvement is usual: mostly papillitis, sometimes optic atrophy.1–3 Three types of macular involvement were described: cystoid macular oedema,1,2,4 geographic atrophy of the macula,1–3 and macular serous detachment.1,2

Macular subretinal neovascularisation has been described in numerous ocular diseases: senile macular degeneration,5 degenerative myopia,5 presumed histoplasmosis syndrome,6–9 acute posterior multifocal placoid pigment epitheliopathy,10 serpiginous choroiditis,11,12 toxocara canis,10,13 chronic uveitis,14,15 coccidiodomycosis,10 drusen of the optic disc,16,17 choroidal naevi,18–21 melanoma and choroidal haemangioma,10,18 choroidal metastasis,10,18 choroidal rupture,22,23 angiod streaks,24,25 pigment epithelial hamartomas,26 fundus flavimaculatus,27 vitelliform macular dystrophy,28 and after argon laser photocoagulation.29,30 These new vessels were also described in systemic diseases: Groenblad-Strandberg syndrome,24 Paget’s disease,25 Ehlers-Danlos disease,10 sickle cell haemoglobinopathies,31–33 sarcoidosis,10 pseudotumour cerebri,34,35 rubella,36–38 Behcet’s disease,39 and Vogt-Koyanagi-Harada syndrome.40

Although macular scars thought to be secondary to disciform macular degeneration41,42 have been observed, we believe there is no published report of active subretinal neovascularisation associated with birdshot retinochoroidopathy. We here report 3 such cases.

CASE REPORTS

CASE 1

In March 1981 we examined a 40-year-old woman with a history of decreased visual acuity and metamorphopsia of the left eye for one month. Her family history was not relevant. There was no evidence of systemic disease, although she had transient hypertension during one pregnancy and was moderately obese. Laboratory studies showed a reduction of the C4 fraction of complement and the presence of antigen HLA B5.

On examination her best corrected visual acuity was RE 6/6 (with −3.75 (−1.25) x 115) and LE 6/9 (with −2.75 (−100) x 20). There was a mild cellular reaction in the anterior chambers. Applanation tonometry was normal. Biomicroscopy disclosed cells diffusely throughout the vitreous of both eyes. There were scattered, discrete, cream coloured hypopigmented subretinal patches in both eyes. Most were less than 1 disc diameter, but in the inferior nasal quadrant some were confluent (Fig. 1). Large choroidal blood vessels appeared normal. There was mild blurring of the disc margins. The hypopigmented patches were less prominent angiographically than

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ophthalmoscopically. During the late phases of the angiogram bilateral leakage of dye from the capillaries of the optic nerve head was evident.

In addition there was a juxtapapillary retinal serous detachment surrounded by deep haemorrhage in the left eye which did not extend into the fovea. A deep greyish lesion, slightly elevated, was located in the centre of the detachment (Fig. 2). A fluorescein angiogram showed an early deep hyperfluorescence, increasing markedly in late phase, located at about 1000 μm from the centre of the foveola (Figs. 3A, B). The Goldmann visual field showed a relative para-central scotoma. Colour testing showed a mild acquired dyschromatopsia. Laser photocoagulation was applied to the neovascular membrane.

CASE 2
A 43-year-old woman had had blurred vision for 3 years. In 1978 she was treated with corticosteroids for vitritis and papillitis. Her medical history was significant in that she had hypertension treated by methyldopa. Her best corrected visual acuity was in January 1980, RE 6/9 (+1-00) and LE 6/6 (+1-00). Biomicroscopy disclosed vitreous cells bilaterally. On fundus examination the typical depigmented lesions extended from the optic disc to the equator. Some patches involved the posterior pole (Fig. 4).

In the right eye there was a juxtapapillary serous detachment, 1 disc diameter in size, which did not extend into the fovea. In the centre of the detachment the angiogram showed an early hyperfluorescence of a deep neovascular membrane extending temporally (Figs. 5A, B) with dye leakage during the study (Fig. 5C).

CASE 3
A 58-year-old woman complained of markedly decreased visual acuity in both eyes over several
years. In May 1981 her best corrected visual acuity was in RE 1/40 and in LE 1/20. Slit-lamp examination disclosed cells in the anterior chamber and in the vitreous. In both eyes the typical subretinal patches of birdshot retinochoroidopathy were present. In her right eye there was a macular scar probably secondary to subretinal new vessels. In the left eye a serous retinal detachment extended from the temporal margins of the disc toward the macula. A haemorrhage was located in the macula (Fig. 6).

In October 1981 the visual acuity in the left eye was unchanged. A fundus examination showed the extension of the serous detachment outlined by a subretinal haemorrhage temporally to the macula (Fig. 7A). On fluorescein angiography a deep early hyperfluorescence underneath the macula was visible (Fig. 7B). This hyperfluorescence increased during the late phases (Fig. 7C). From the margin of the optic nerve a glial proliferation was extending towards the macula. These angiographic features implied the presence of subretinal neovascular tissue.

**Discussion**

The occurrence of subretinal new vessels in uveitis has been recorded at a cicatricial stage of focal choroiditis.

In "presumed ocular histoplasmosis syndrome" the new vessels usually arise from the atrophic scars (histospots). In toxoplasmosis the subretinal new vessels originate from the retinal choroiditis focus. Such cases have been published of patients with infection by toxocara canis, syphilis, and tuberculosis. In all these cases the retinal pigment epithelium and Bruch's membrane were partially or totally destroyed at the site of the atrophic scars. Thus the new vessels of choroidal origin had access to the subepithelial or subretinal space of the macular area.
Chronic intraocular inflammation may be associated with a diffuse and extensive depigmentation of the retinal pigment epithelium. In diffuse and nonspecific chronic uveitis the occurrence of subretinal new vessels is uncommon. Few cases have been published—one with peripapillary neovascularisation by Schwartz, and 2 by Augsburger (one with peripapillary neovascularisation, one with bilateral macular new vessels). These authors reported pigment epithelium disturbances of the posterior pole which evoked atrophic scars. They believe that chronic ocular inflammation stimulated the proliferation of subretinal new vessels located near the atrophic patches involving the pigment epithelium and the choriocapillaris.

The characteristic feature of the birdshot retinochoroidopathy is the existence of yellow-white or faintly orange patches of depigmentation. These flat patches are usually symmetrically distributed in both eyes, scattered in the postequatorial part of the fundus, and often more numerous in the area nasal to the optic disc than elsewhere. There is no evidence of hyperpigmentation within the lesion or at their margins. There is no slit-lamp evidence of thinning either of the retina or of the choroid in the area of the patches. Large choroidal blood vessels are usually observed within the lesion in white light but the overlying retinal vessels appear normal.

During the early phase of angiography the patches show no evidence of alteration in the normal background of choroidal fluorescence. There is no evidence either of early hypofluorescent areas to suggest choriocapillaris atrophy, or early hyperfluorescent areas that might be expected if there were depigmentation of the pigment epithelium or loss of function of the pigment epithelial barrier. This
would be in accord with the suggestion by Gass that the pallor of the lesions was due to loss of pigment from choroidal melanocytes. Such an appearance could also be produced by the presence of abnormal deposits beneath the pigment epithelium more evident in white light. Such deposits may induce neovascularisation or alter Bruch's membrane so that a barrier to new vessel membrane growth is destroyed.

The peripapillary location of the new vessels in all 3 cases may suggest that inflammation of the nerve head and of the peripapillary region is of importance in the genesis of neovascularisation. There is no evidence that inflammation has little if any effect on the retinal pigment epithelium, but it has on Bruch's membrane. Recently it has been suggested that although the occurrence of birdshot retinochoroidopathy is genetically predisposed, retinal autoimmunity plays a role in the manifestation of the disease.

Serous detachment and haemorrhages have been
Case 3. Red-free photograph: retinal serous detachment temporal to the optic nerve outlined by a haemorrhage; scattered birdshot patches.

Fig. 6

Fig. 7A

Fig. 7B

Fig. 7C

Fig. 7 Case 3. Five months later: extension of the serous detachment on the other side of the foveola on red-free photograph (A). Early hyperfluorescence in the macula outlined by a deep haemorrhage (B). Late phase: leakage of dye in the macula and fibrous tissue extending from the optic disc (C).

reported in birdshot retinochoroidopathy, but in no case described were subretinal new vessels identified. The decreased visual acuity in this disease, with many manifestations and a broad spectrum, may be due to optic atrophy or cystoid macular oedema. Subretinal neovascularisation is another possible sight-threatening complication.

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References


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