Fluorescence in Best’s vitelliform dystrophy, lipofuscin, and fundus flavimaculatus

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SUMMARY Control photographs, with the Baird Atomic B4 and B5 filters in place prior to fluorescein injection, show exposure of the film corresponding to (1) the small yellow vitelliform lesions at the edge of a disrupted disc, (2) the pseudohypopyon in a vitelliform cyst, (3) orange lipofuscin overlying a malignant melanoma, and (4) some of the flecks in a case of fundus flavimaculatus. Because of transmission overlap between the filters, the relative contribution of reflected light and true autofluorescence is difficult to quantitate. Reflectile structures such as the optic nerve or a white scar were essentially unexposed, but minimal fundus detail was seen. Some parallels exist between lipofuscin and the content of a disrupted vitelliform lesion.

‘Autofluorescence’, primary, or natural fluorescence, is a physical property in which a light-absorbing substance (fluorophore) re-emits a longer wavelength of light. In fundus fluorescein angiography this optical phenomenon must be differentiated from two types of pseudofluorescence. The first occurs when light reflected from highly reflecting fundus structures excites residual fluorescein in the aqueous and vitreous humour. This excited residual fluorescein in the later stages of angiography may cause a photographic image (Machemer et al., 1970; Archer, 1972). The second type, ‘pseudoautofluorescence,’ is a reflectile phenomenon in which a faint fundus image, resulting from transmission overlap of the filter pairs, is discernible in control photographs.

Autofluorescence of superficial optic nerve head drusen has been a helpful fluorescein angiographic finding in distinguishing drusen from other causes of disc swelling (Lorentzen, 1966; Sanders and Fytyche, 1967). Many clinicians have noted fluorescence of extensive hard exudate. However, this phenomenon has not been reported in other clinical entities. This report concerns the observation of preinjection fluorescence of (1) the disrupted egg yolk deposits in Best’s vitelliform macular dystrophy, (2) orange lipofuscin pigment over a choroidal melanoma, and (3) some of the flecks in a case of fundus flavimaculatus.

Patients and methods

Three members of a family had multifocal, macular

and extramacular, Best’s vitelliform dystrophy. The diagnosis of Best’s dystrophy seemed secure on the grounds of (1) occurrence in a parent and 2 children, (2) characteristic macular lesions, and (3) electrooculogram light-peak/dark-trough ratio between 1.0 and 1.12 in all eyes (normal greater than 1.85).

Fluorescein angiography was performed in the standard manner. Control photographs were taken before fluorescein injection. The filter pair used was the Baird Atomic B4 exciter (less than 2% of the excited light is transmitted above 5000Å) and the B5 barrier (less than 2% is transmitted below 4950Å and 0.1% below 4900Å). Actual transmission curves of the filters used are graphed in Figs. 1 and 2.

Six other patients with Best’s vitelliform dystrophy and fluorescein angiography were reviewed. Because of the colour of the orange yellow egg yolk in Best’s dystrophy and the orange lipofuscin pigment seen overlying a choroidal melanoma, colour photographs were reviewed on the 30 most recent cases seen here with the histologically proved diagnosis of malignant melanoma of the choroid. Only 1 case had prominent orange pigment. Fluorescein angiography was available in 6 of 12 patients with the diagnosis of fundus flavimaculatus.

Results

All yellow disrupted egg yolk spots demonstrated preinjection fluorescence (Figs. 3a, b, c). This fluorescence occurred in the pseudohypopyon as well as in the circinate distributed yellowish white deposits of the remaining ‘scrambled egg’ lesions. There was relative fading of the optical phenomenon as the arteriole phase began. The yellow deposit of
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B4 Exciter filter transmission curve

Fig. 1

the pseudohypopyon and the extramacular lesion blocked fluorescence during the fluorescein transit. The circinate yellow deposits acted unpredictably, with some blocking and some transmitting background fluorescence, but most stained. Intact egg yolks, diffuse chorioretinal atrophic lesions, or end-stage fibrous scars did not fluoresce. The orange pigment overlying the choroidal melanoma fluoresced in control pictures (Figs. 4a, b). The pigment blocked fluorescence in the later stages of angiography. One of the 6 patients with fundus flavimaculatus demonstrated preinjection fluorescence (Figs. 5a, b; 6a, b). During the dye transit, these flecks showed a retinal pigment epithelial transmission defect, late photographs demonstrated staining of these flecks.

Discussion

The cases presented demonstrate fluorescence of (1) the small multiple vitelliform deposits at the margin of the larger scrambled atrophic lesion, (2) the pseudohypopyon of a vitelliform cyst, (3) the orange lipofuscin overlying malignant melanoma of the choroid, and (4) some of the flecks in fundus flavimaculatus.

Autofluorescence is a re-emission phenomenon—a shorter wavelength of light is absorbed and a longer one emitted. It is not a reflective phenomenon like pseudoautofluorescence—a faint fundus image discernible in control photographs resulting from transmission overlap of the filter pairs. Likewise, pseudofluorescence is excitation of fluorescein within the ocular media from light reflected from reflectile fundus structures such as a white scar.

Indeed, autofluorescence is a common biological phenomenon. Nearly all proteins excited in the 2500-2800Å region will fluoresce because of the presence of tryptophan, tyrosine, and phenylalanine. Substances that normally have strong autofluorescence include (Pearse, 1962) collagen, elastic fibres, protein bound NADH<sub>2</sub>, vitamin A, lipofuscin, and porphyrin. Not unexpectedly, the cornea, nuclear sclerotic lens and intraocular fluids (Krill, 1972) have autofluorescence. However, with conventional angiography this is either minimal or non-detectable.
Despite the efficient separation between the transmission characteristic of the exciter B4 and the barrier B5 filters, there is nonetheless some low-intensity overlap in transmission, especially of longer wavelengths. Therefore the possibility remains that the fluorescing substance in Best's vitelliform dystrophy, lipofuscin granules and some of the flecks in fundus flavimaculatus, is not autofluorescent but rather reflecting enough light to expose the film. Indeed, minimal fundus detail can be seen in most

Fig. 3a, b, c Preinjection fluorescence of a variety of macular and extramacular lesions in Best's vitelliform dystrophy

Fig. 4a Choroidal malignant melanoma temporal to the macula in the left eye. The tumour surface is peppered with orange pigment. A confluent patch of orange pigment is present infranasally (arrow)

Fig. 4b Preinjection photographs with the Baird Atomic 4 and 5 filters in place demonstrate fluorescence of the orange pigment (arrow)
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of the mounted pictures. This visibility attests to the fact that the filter combination is not ideal. However, those structures that would be expected to reflect light such as the white scar and the optic disc were not visible on the control photographs at the same exposure in which the fluorescing substance was vividly visible. A more perfect filter pair would help resolve the relative contribution of reflected light and autofluorescence. Indeed quantitative absorption and emission spectra of these substances might identify them.

Whether there is any common substance between Best’s vitelliform macular dystrophy, fundus flavimaculatus, and lipofuscin is pure speculation. Nonetheless, there are a few interesting common points between Best’s dystrophy and lipofuscin as summarised in Table 1.

1. Lipofuscin is orange. *a.* The vitelliform disc is an orange to yellow colour.

2. Lipofuscin granules are most prominent in the macular region. *a.* The vitelliform disc is almost always macula centred.

3. Lipofuscin increases with age at the posterior pole (Streeter, 1961; Friedman and Tso, 1968). *a.* Vitelliform lesions often become more visible or symptomatic with age (Deutman, 1971) (during its early evolution; after disruption they become less evident).

4. Lipofuscin autofluoresces, with its maximum at 4300 to 4500Å. *a.* The disrupted vitelliform deposit and the lipofuscin overlying a malignant melanoma fluoresce under the same photographic setting.


6. Lipofuscin occurs predominantly within retinal pigment epithelial cells. *a.* Best’s dystrophy is thought to be basically a disturbance of the retinal pigment epithelial cell.
Table 1

<table>
<thead>
<tr>
<th>Colour</th>
<th>Best's vitelliform disc</th>
<th>Lipofuscin</th>
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<tbody>
<tr>
<td>Location</td>
<td>Macula centred</td>
<td>Greatest concentration in macula</td>
</tr>
<tr>
<td>Change with age</td>
<td>More visible, symptomatic in early stages (fading late)</td>
<td>Increasing density</td>
</tr>
<tr>
<td>Preinjection fluorescence</td>
<td>4000-4500Å</td>
<td>4300-4500Å</td>
</tr>
<tr>
<td>Fluorescein characteristic</td>
<td>Blocks transmission</td>
<td>Blocks transmission</td>
</tr>
<tr>
<td>Cell involvement</td>
<td>Probably retinal pigment epithelium</td>
<td>Retinal pigment epithelium and macrophages</td>
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<tr>
<td>Pathological occurrence</td>
<td></td>
<td>1. Vitamin-E-deficient animals (Mason and Hartsough, 1951; Hayes, 1974)</td>
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<td></td>
<td></td>
<td>2. Under and within degenerative retina overlying malignant tumours</td>
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7. a. Acid-fast yellowish-brown autofluorescent pigments (histochemical characteristics of lipofuscin) were observed to develop in vitamin-E-deficient rats, mink, pigs (Mason and Hartsough, 1951), and monkeys (Hayes, 1974). b. Lipofuscin over a choroidal melanoma is located within proliferating retinal pigment epithelial cells and in macrophages under and within degenerated retina. c. In neuronal ceroid-lipofuscinosis there is early loss of rods and cones, with degeneration of melanin-containing epithelial cells. Characteristic autofluorescent lipopigment granules are found in ganglion, Müller, and photoreceptor cells (Goebel et al., 1977).

At the same time one must bear in mind that lipofuscin is not a single substance but rather a lysosomal oxidative product of lipid or lipoprotein from cell breakdown.

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References


