COMMUNICATIONS

TOXOPLASMOSIS IN THE ADULT*

BY

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Toxoplasmosis as a congenital infection is now a well-established clinical entity, but our knowledge of toxoplasmosis acquired in childhood or adult life is much less complete. It is known, however, that such infections may occur, and several cases without ocular involvement have been reported (Pinkerton and Weinman, 1940; Pinkerton and Henderson, 1941; Guimarães, 1943; Syvertson and Slavin, 1946; Kean and Grocott, 1947; Noetzel, 1951), and Sabin (1942) described an acquired form of the disease in older children. Most of the cases of acquired toxoplasmosis in man have not shown ocular involvement, and Sabin (1950) and Sabin and others (1952), have emphasized the fact that there is as yet no satisfactory evidence that the organism can cause a chorio-retinitis through a post-nataly acquired infection. Although he did not deny its possible occurrence, Sabin (1950) pointed out that the fully developed and post-natal nervous tissues are unquestionably less susceptible to the pathogenic effects of toxoplasma than are those of the embryonic nervous system.

Nevertheless, there are in the literature a few cases of toxoplasmosis with chorio-retinitis where acquired infection in childhood or adult life has been suspected. Six cases of chorio-retinitis associated with toxoplasma were reported by Vail, Strong, and Stephenson (1943) and, in their view, one patient aged 16 years, undoubtedly had an acquired infection. Magnusson (1951) reported seven cases of adult infection (two of whom had been infected during their work in the bacteriological laboratory); one of the patients had acute chorio-retinitis and the serological tests supported the diagnosis of toxoplasmosis. Wising (1952) described the case of a female aged 31 years, who presented with lymphadenopathy, pyrexia, and an acute juxtamacular chorio-retinitis in the left eye; the clinical diagnosis of toxoplasmosis was confirmed by strongly positive dye and complement-fixation tests. Rieger (1951, 1952), reported fourteen cases of central exudative retinitis in adults, which, on the basis of skin and serum tests, were considered to be due to acquired toxoplasmosis.

It is not yet established whether a state of chronic toxoplasmosis can exist in man, as it may in animals. Although the fact that mothers who give birth to toxoplasma-infected diseased infants do not themselves show clinical

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evidence of the disease proves that the condition may occur in a latent or sub-clinical form, the normality of all subsequent pregnancies argues against a prolonged chronic infection. On the other hand, a case reported by Bohn and Koch (1951), suggests that toxoplasma may persist in the body in a latent form for many years. This patient, a female aged 22 years, developed acute chorio-retinitis, the dye tests were positive, and cerebral calcification was present; this, according to the studies of Johnson, Fried, Broaddus, and Lamfrom (1946), probably indicates a congenital infection or one acquired in early childhood. Furthermore, an account of an interesting family showing a syndrome of chronic meningo-encephalitis, chorio-retinitis, rash, nerve-deafness, and eosinophilia due to toxoplasmosis, was reported by Campbell and Clifton (1950); it was apparent that the infection had remained in a subacute form in three, and possibly four, members of the family for many years. Kean and Grocott (1947) found an aggregation of parasites, probably toxoplasma, in the cardiac muscle of a healthy Negro who committed suicide, and Plaut (1946) also demonstrated toxoplasma in the myocardium in a child and in an adult. It would appear possible, therefore, that an active chorio-retinitis in an adult may be due not only to a recently acquired infection, but also to an exacerbation of an old infection either post-natally or congenitally acquired. That such infection may be more common than is indicated by the few published cases, is suggested by the recent findings of Wilder (1952a, b), who claimed to have demonstrated organisms morphologically indistinguishable from toxoplasma in granulomatous chorio-retinal lesions in 53 eyes of adults; the protozoa were found in 30 out of 131 eyes with the pathological diagnosis of tuberculosis or possible tuberculosis. The lesions in which the organisms were found were very similar and consisted of well-demarcated necrotic areas involving the retina and choroid and frequently the sclera. They were situated posteriorly, equatorially, or peripherally, and occasionally more than one lesion was present in a single eye. Sabin and others (1952), however, believe that it is better to reserve judgment on this interesting communication until the results of serological tests on the patients from whom the eyes were removed are available, or until such time as toxoplasma organisms from similarly affected enucleated eyes, have been demonstrated by animal inoculation.

Further evidence on this subject is presented in this paper, wherein the clinical, serological, and histological findings in a case of chorio-retinitis in an adult are reported, together with a brief account of the histological examination of 32 cases of granulomatous uveitis.

Case Report

A male, aged 40, first came under observation in August, 1947, giving an ocular history that the vision in his left eye had been defective "as long as he could remember." Presumably this dated from youth. Ten years previously (1937) the vision in this eye had deteriorated further and at that time bilateral antrostomies had been performed. During the 10 weeks previous to seeking advice he had noticed that the vision in the left
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eye had again become worse. The general medical history contained nothing significant; but there was a history of tuberculosis in the family.

Examination.—The right eye was in every respect normal—vision 6/5. In the left eye the anterior segment was normal; there was a considerable amount of fine vitreous opacities, and from the upper temporal margin of the disc there extended a large patch of deep choroiditis showing an inflammation of the acute granulomatous type superimposed on an old lesion of long standing. The remainder of the fundus was normal; there was an absolute field defect corresponding to the chorio-retinal lesion; vision was 6/18.

Diagnosis.—The patient was hospitalized and overhauled with a view to establishing the diagnosis. In general he appeared to be extremely healthy. Bacteriologically the only findings which could be interpreted as possibly pathological were a coagulase-positive staphylococcus in the upper respiratory tract and a streptococcus haemolyticus, type 3, in the upper respiratory passages and bowel. There was a strongly positive Mantoux reaction to human tuberculin and a weakly positive reaction to bovine tuberculin; but all other serological tests were negative. Toxoplasma tests were not carried out at this stage. There was no dental infection and x rays of the chest were negative.

The appearance of the eye did not suggest the likelihood of the infection being streptococcal, and, in view of its clinical characteristics, the family history, and the strongly positive Mantoux reaction with the differentiation between the response to human and bovine tuberculin, the case was provisionally labelled tuberculous.

Therapy.—After the acute stage of choroiditis had passed, the patient was given a course of tuberculin injections, starting with minute doses.

Later Developments.—In October, 1947, the lesion was quiet, leaving an atrophic area spreading out widely from the disc: there was still a slight vitreous haze: vision 6/18.

The eye remained quiet until August, 1949, when the patient returned with a history that the vision in the left eye had deteriorated suddenly 5 days previously. On examination the anterior segment was again found to be normal, a dense vitreous haze obscured all view of the fundus, central vision had gone, and hand-movements only were apparent in the peripheral field. With routine treatment the posterior uveitis gradually settled down, and by February, 1950, the vitreous was relatively clear and an immense area of atrophic scarring was seen to occupy the posterior pole involving the macula and embracing the disc, associated with optic atrophy. An extensive relapse of a necrotizing choroiditis had obviously occurred.

The eye remained quiet until September, 1950, when there was again a sudden deterioration of vision, for the first time associated with pain. On examination the ocular tension was found to be raised, the cornea was oedematous with its posterior surface bespattered with keratic precipitates, the iris was heavily vascularized, no view was obtainable behind the pupil, and all perception of light had gone. The clinical picture suggested the presence of a vascular thrombosis of considerable extent. On hospitalization and with intensive local treatment the tension fell and the acute symptoms subsided: but since a considerable degree of discomfort and photophobia remained, the eye was excised in November, 1950. From that time onwards the patient has remained in excellent health.

Summary.—Recurrent posterior uveitis of obscure aetiology, clinically of the granulomatous type, persistently recurring, resulting in loss of perception of light and eventual destruction of the eye.

Pathological Findings

In November, 1950, the globe was opened horizontally to reveal a chorio-retinal ulceration in the posterior temporal region. Horizontal celloidin sections were cut and stained with haematoxylin and eosin, Van Gieson’s stain, and Ziehl-Neelsen’s stain.

Sections.—There was a mild interstitial inflammatory infiltration of the corneal stroma and small foci of lymphocytes could be seen within the corneal epithelium. Massive
keratic precipitates were present on the corneal endothelium and the filtration angles were partially occluded by aggregations of inflammatory cells; posterior synechiae were present. Lens normal.

Throughout the uveal tract there was a chronic inflammatory infiltration, severe in the iris, in which nodular foci were present at the pupillary margin, moderate and diffuse in the ciliary body and anterior choroid—a delicate cyclitic membrane was present—and severe in the posterior region where there was a well-demarcated area of necrosis on the temporal side. In this region the retina and choroid were fused, being divided only by a strip of pigment epithelium and remnants of Bruch’s membrane (Figs 1, 2, and 3); in the centre of this lesion the retina was necrotic and had completely lost its normal architecture. There was a mild chronic inflammatory infiltration in the underlying sclera.

Elsewhere the retina showed severe perivascular cuffing and nodular proliferations of endothelial cells, especially around the retinal veins and on the internal limiting membrane.

Fig. 1.—Section of globe. On the temporal side of the disc there is a well-demarcated area of granulomatous chorio-retinitis. See Fig. 2.

Fig. 2.—High-power view of area of chorio-retinitis. Retina and choroid are fused, being separated only by a layer of pigment epithelium and remnants of Bruch’s membrane. The lesion is granulomatous and centrally necrotic. The underlying sclera shows a mild chronic inflammation. Haematoxylin and eosin × 22.
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Fig. 3.—Severe chronic inflammatory infiltration of iris with nodular aggregations at pupillary margin. Haematoxylin and eosin × 44.

Some of the smaller retinal vessels had become obliterated by the exuberant tissue proliferation. Scattered eosinophils were present in and around the vessels, but these cells were not prominent; a few giant cells could be seen in the area of retinal necrosis and in association with the endothelial proliferations elsewhere in the retina, but were not present in the choroid. There was cellular vitreous exudate, particularly dense over the ciliary processes; the disc was oedematous and showed a severe inflammatory infiltration. Ziehl-Neelsen staining revealed no tubercle bacilli.

The histological diagnosis at this time was chronic granulomatous chorio-uveitis, probably tuberculoid in origin. A Middlebrook-Dubos test and agglutination tests for Brucellosis were carried out with negative results.

In August, 1952, in view of the reported findings of Wilder (1952a, b) the sections were re-examined 21 months after excision.*

Sections.—Serial sections were cut throughout the necrotic chorio-retinal lesion, periodic acid-Schiff and Heidenhain’s iron haematoxylin being used in addition.

Many crescentic structures morphologically indistinguishable from toxoplasma were found in the necrotic area of the retina and in the nodular lesions of the iris (Figs 4, 5,

Fig. 4.—V-shaped pair of crescentic structures indistinguishable from toxoplasma. Nuclei localized at adjacent poles. (Authors’ case). Haematoxylin and eosin × 2560.

Fig. 5.—Two separate crescentic forms. Lower organism shows a central nucleus and is identical with toxoplasma (Authors’ case). Heidenhain’s haematoxylin and Masson stain ×2540.

* Dr. Helenor Wilder herself selected the slide from this case as a probable example of toxoplasmosis.
6, and 7). In some of the sections over twenty scattered crescentic forms were found, while in others only three or four were observed. Some occurred in pairs, either parallel or in a V-shape, the nuclei being located at the tip of the V. The majority, however, were isolated, the nuclei being situated either at the middle or at one end, and granules were sometimes distinguishable in the cytoplasm. A few of the forms were intracellular, but no typical pseudo-cysts or cysts containing toxoplasma were seen. With haematoxylin and eosin the cytoplasm stained faintly blue and the nuclei dark blue. P.A.S. stain was not sufficiently selective to be of use; Heidenhain's iron haematoxylin was of the greatest value in differentiating the organisms (Smith, 1953). These findings indicated a probable toxoplasmosis, and serological tests were carried out in August, 1952, that is, 2 years after the last relapse of the uveitis. These tests were repeated in October, 1952, and in January, 1953.

Serological Tests (performed by Professor Beattie, Sheffield University):

A. Cytoplasm Modifying Dye Test.
   1. (18.8.52) Positive up to 1:160.
   2. (18.10.52) Positive up to 1:68.
   3. (26.1.53) Positive up to 1:32.

B. Complement-fixation Test.
   1. (18.8.52) Positive 1:2.
   2. (26.1.53) Negative.

C. Rabbit Skin Test.
   1. (18.10.52) Serum neutralized more than ten and less than one hundred skin-test units.
   2. (26.1.53) Ten skin-test units neutralized.

D. Dermal Sensitivity Test to Toxoplasmin.
   1. (10.10.52) Strongly positive.

Discussion

In the interpretation of the cytoplasm modifying dye test there is not yet complete agreement on the levels of the positive titres which may be regarded as significant, and to some extent this is due to the variation in the range of positive results in the healthy populations of different countries. According to Sabin and others (1952), however, the titres do not exceed 1:64 in the vast majority of the normal population, and this figure is in accord with the findings of Beverley and Beattie (1952) in Great Britain:

Fig. 6.—Nucleated crescentic form in choroidal granulation tissue (Authors' case). Haematoxylin and eosin × 1600.

Fig. 7.—Choroidal granulation tissue containing crescent forms, one showing central nucleus (Authors' case). Haematoxylin and eosin × 1600.
Thus the figure of 1:160 obtained in our case, is of possible significance, and becomes even more striking when one considers the rapidly falling titre, which, if due to the removal of the antigenic focus 21 months previously, would suggest a much higher level at the time of the acute phase of the uveitis. Indeed, it is reasonable to suppose that this patient’s left eye was the only remaining tissue to have contained viable protozoa; Weinman (1943) has shown that there is a differential visceral immunity in toxoplasmosis with immunity marked in the lung and feeble in the brain, a phenomenon which has been attributed to a lesser permeability of the cerebral capillaries to antibodies as compared with capillaries elsewhere (Friedemann, 1942). It is possible that the organisms are similarly shielded within the retinal tissues, and that, in the particular case here reported, they had survived only within the excised eye.

With regard to the complement-fixation tests, it has been found that titres as low as 1:2 can be specific, in that such sera have also invariably contained significant amounts of cytoplasm-modifying antibody (Sabin and others, 1952). The same authors have established that the complement-fixing antibody appears later and disappears much earlier than the cytoplasm-modifying antibody, so that the findings in our case (a conversion of a positive titre of 1:2 to a negative result in 5½ months) is in accord with a recovery from an acute infection. The rabbit skin tests and the dermal sensitivity tests are confirmatory.

We therefore conclude that the serological results are to be regarded as significant, and that, when these are taken in conjunction with the histological findings, there can be little doubt that the granulomatous chorioretinitis was toxoplasmic in origin. The existence of crescentic forms in the necrotic inflammatory tissue and the absence of pseudo-cysts would appear to indicate an acute phase of the infection, as pointed out by Weinman (1952), while the clinical history of defective vision “as long as he could remember” would suggest a congenital infection or an onset in early childhood. On the other hand, according to Johnson, Fried, Broadus and Lamfrom (1946), the absence of cerebral calcification indicates an onset after the age of 15 years. The question whether this was a case of congenital infection or of infection acquired post-natally in childhood must, therefore, remain in doubt, but the evidence is in favour of a chronic infection remaining latent in the ocular tissues for many years and recrudescing to give rise to an acute chorioretinitis in middle age. Thus the findings in this case support in a rather striking way the opinion of Wilder (1952b) that, in adults, granulomatous chorioretinitis clinically and histologically resembling tuberculosis may be due to toxoplasmosis. Wilder does not state the number of eyes examined, but the fact that the protozoa were demonstrated in 53 cases

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<th>Under 10 years</th>
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<td>1 : 16 or over.</td>
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suggests that toxoplasmosis may be a not uncommon cause of chorio-retinitis in adults. In order to ascertain whether a similar high incidence is to be expected in Great Britain we have examined sections from 32 cases of granulomatous uveitis reported at this Institute since 1948.

None of the 32 cases had the localized type of necrotic lesion described by Wilder as typical of toxoplasmosis. Nevertheless, in about half of the cases (17) a few structures were found resembling, to a greater or lesser extent, crescentic toxoplasma forms, but their appearance was not sufficiently typical to identify them as protozoa, nor were their isolated disposition and sparse distribution (never more than five in one case) characteristic of toxoplasmosis. That similar crescentic forms were also found in cases of post-traumatic granulomatous uveitis (Fig. 8), in which toxoplasmosis was not in question, casts further doubt upon their nature. Furthermore, a solitary, isolated crescentic form with a typical nucleus was demonstrated in a case of Behcet’s syndrome (Fig. 9). It was apparent that degenerate cells undergoing karyolysis and karyorrhexis may often very closely resemble toxoplasma pseudo-cysts. In one case in which crescentic structures were found serological tests were carried out (dye test 1:8; complement-fixation test negative).

Thus from this relatively small group of patients it may be concluded that in cases of granulomatous uveitis neither resembling tuberculosis nor readily admitting a diagnosis of toxoplasmosis, histological structures morphologically similar to toxoplasma (probably consisting of cell debris or nuclear debris) may be found.

Further investigation on this important subject is clearly necessary, but in view of the difficulty in interpreting suspicious structures in histological material, we feel that great caution should be exercised before labelling them
as toxoplasma. Before regarding the diagnosis as even probable, it is necessary, in our opinion, to have observed quite typical histological forms—crescents in pairs being particularly significant—and to have demonstrated positive serological tests.

When sufficient examinations fulfilling the above criteria have accumulated, it may well be that Wilder's suggestion that toxoplasmosis is a not uncommon cause of chorio-retinitis in adults may be generally confirmed. Meanwhile it should be emphasized that there is as yet no certain evidence to show that the case here reported is other than a rarity.

**Summary**

(1) A case of recurrent granulomatous chorio-retinitis in an adult is reported, and clinical, histological, and serological evidence is given for regarding the condition as due to toxoplasmosis. It is suggested that the disease was either congenital or acquired in childhood, and that the active chorio-retinitis in adult life was due to a recrudescence of the early infection.

(2) When sections of 32 cases of granulomatous uveitis were examined, about half showed structures resembling toxoplasma forms. However, in some of the cases in which these were found, which included such conditions as post-traumatic uveitis, and Behcet's syndrome, the diagnosis of toxoplasmosis would have been highly improbable. It is, therefore, concluded that these histological appearances probably result from cellular debris or nuclear fragmentation, and that, in the absence of confirmatory serological tests, such findings have a very limited value.

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