Experimental autoimmune uveoretinitis in the RCS rat: the influence of photoreceptor degeneration on disease expression

E G Atkinson, C Edelsten, Eva Kasp, D C Dumonde

Abstract
S-antigen induced experimental autoimmune uveoretinitis (EAU) was produced in the Royal College of Surgeons (RCS) strain of rat which develops a photoreceptor dystrophy within 2 weeks of birth. Animals were sensitised at 60, 90, and 105 days of age: all animals developed disease, but onset was significantly delayed in older (105 day) animals compared with those aged 60 days at sensitisation (p<0.003). Disease was characterised by the early development of complete serous retinal detachment which resolved in a few days: the prevalence of retinal detachment was increased to 80% in dystrophic animals compared with 10% in the congenic, non-dystrophic controls (p<0.001). Anterior uveitis was seen in 17/30 control strain eyes, but in none of 30 dystrophic eyes (p<0.001). Genetically determined photoreceptor and retinal pigment epithelium dysfunction in the RCS rat, which may involve the local accumulation of altered S-antigen, predisposes the dystrophic strain to display an acute retinal detachment in the early stages of EAU. This phenomenon illustrates how biochemical dysfunction of a target organ may influence susceptibility, form, and severity of an experimental autoimmune disease.

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Experimental autoimmune uveoretinitis (EAU) is a model of human uveoretinal inflammation, produced by sensitisation of susceptible animal species to discrete retinal proteins, and results in photoreceptor necrosis. Retinal S-antigen, which is confined to the photoreceptors and pineal gland, was the first purified protein demonstrated to produce EAU, and has been the most intensively studied.1 Autosensitisation to retinal proteins is considered possible because of their anatomical sequestration from the developing immune system. When immunocompetent animals are then sensitised to S-antigen, T helper cells and, later, T cytotoxic/suppressor cells enter the photoreceptor layer and are presumed to produce the characteristic damage.2 Disruption of the blood-retinal barrier and destruction of photoreceptors promote further access to and recognition of the target antigens by the sensitised immune system. On this basis, continued interaction between the immune system and components of the photoreceptor layer depends on the continuing availability of photoreceptor antigens to the sensitised immune system. Indeed, when the normal turnover of S-antigen at the photoreceptor outer segments is disrupted, as is seen after retinal light damage or dark adaptation, the severity and chronicity of S-antigen induced uveoretinitis is reduced.3

Against this background it seemed likely that susceptibility to and severity of EAU could well be affected by coexisting photoreceptor pathology. The Royal College of Surgeons (RCS) strain of rat develops an inherited photoreceptor dystrophy: its abnormal retinal pigment epithelium is unable to phagocytose effectively the photoreceptor outer segment membranes. The photoreceptors start to degenerate within 2 weeks of birth, and cellular debris accumulates spontaneously in the subretinal space: by 2 months of age, the outer nuclear layer is reduced to a thickness of two nuclei. The subretinal debris begins to disappear by 3 months of age, but some may persist for up to 18 months. Eventually, the inner nuclear layer rests on the pigment epithelium, the retinal pigment epithelium (RPE) becomes vascularised and there is gliosis within the remaining retina.4,5 The black-eyed strain of RCS rat has a pigmented fundus that allows funduscopic and angiographic assessment to be combined.6 We considered that a comparative study of EAU in congenic, non-dystrophic rats of the same strain would enable evaluation of the influence of photoreceptor dystrophy on disease expression. Accordingly we have studied the form and severity of EAU in RCS dystrophic rats, sensitised at different ages, using congenic, non-dystrophic pigmented rats as controls, and characterising the uveoretinitis by funduscopic, angiographic, and histopathological criteria.

Materials and methods
Fifteen dystrophic and 15 control strain RCS rats were sensitised with 50 μg of retinal S-antigen in adjuvant so as to induce EAU. They were examined initially at 2–3 day intervals from 14–45 days after sensitisation, and then followed-up until day 81: fundus photography and fluorescein angiography were used to document changes in the clinical signs. The animals were then killed on day 86 post sensitisation, and the eyes removed for histological examination. Initial experiments9 had suggested that the time of onset of disease in dystrophic rats depended on age at sensitisation, and that no such relationship was found in the control strain. Accordingly in this study the dystrophic animals were sensitised at three different ages: 60 days (five rats), 85–90 days (seven rats) and 105 days (three rats). Non-dystrophic controls were sensitised at 60 days (four rats) and 85–90 days (11 rats). Experiments consisted of documenting the
ophthalmoscopic and histopathological features of uveoretinitis in dystrophic and control strains at early, intermediate, and late stages of disease; and of examining relationships between the age of onset of EAU and age of rats at sensitisation. Strain differences in the prevalence of EAU or of retinal detachment were evaluated by χ² analysis and relationships between time of onset of EAU and age of rats at sensitisation were examined by the Wilcoxon rank sum test.

INDUCTION OF EAU IN RCS RATS
RCS, black-eyed dystrophic rats (rdy/rdy p') were obtained from the Central London Polytechnic breeding colony, together with 15 control strain rats (rdy/p'). They weighed 100-150 g and were maintained under non-specific pathogen free conditions. Animals were sensitised in the hind foot pads with 50 µg of S-antigen in Freund's complete adjuvant (FCA: Sigma) in an injection volume of 100 µl. Bovine S-antigen, prepared by previously published methods, was dissolved in phosphate buffered saline (PBS) to 0.5 mg/ml and emulsified with an equal volume of FCA fortified with heat-killed Mycobacterium tuberculosis (3 mg/ml: MAFF, Weybridge, England). Simultaneously, 10⁴ heat inactivated Bacillus pertussis organisms were given intraperitoneally in 0.25 ml PBS.

CLINICAL AND ANGIOGRAPHIC ASSESSMENT OF EAU
Animals were examined by slit-lamp and indirect ophthalmoscopy, after mydriasis with 1% tropicamide (Smith & Nephew). The presence of the following clinical features were recorded separately on each occasion: anterior uveitis, retinal detachment, disc oedema, retinal venous dilatation and periphlebitis, retinal infiltrates, and abnormalities of the retinal pigment epithelium. Selected features were recorded by fundus photography. Fluorescein angiography was performed under Thalamonal anaesthesia, 0-05 ml/kg (Janssen, Oxford). Sodium fluorescein (4%; 0.1 ml) was injected via the tail vein. Angiograms were recorded using a Zeiss fundus camera using Ilford FP4 film.

HISTOLOGICAL PROCESSING
At termination of the experiment, animals were killed by cardiac puncture under ether anaesthesia. Eyes were removed, fixed in 4% glutaraldehyde (EM grade, Emscope Laboratories) for 12 hours, then 10% formaldehyde-saline (BDH); followed by vacuum wax embedding and haematoxylin and eosin staining.

Results

PREVALENCE AND COURSE OF CLINICAL DISEASE IN RCS RATS
All animals developed EAU (Fig 1). The dystrophic strain tended to develop disease earlier, and resolve quicker. At day 18 there were significantly more dystrophic rats with disease than controls (16/30 eyes vs 8/30 eyes: χ²=4.44, p<0.05), but by day 58 there were significantly fewer dystrophic animals with disease compared with controls (12/30 vs 24/30, p<0.01) (Fig 2). However, there was no significant difference in the time of disease onset between the dystrophic rats [median 18 days (range 15-46)], and controls [23-5 days (15-55)], or the length of active disease between the two groups: 24 days (3-51) in dystrophic strain vs 29 days (8-61) in control strain. Four non-immunised control strain rats and three non-immunised dystrophic rats did not develop uveitis.

When the animals were grouped according to age at immunisation (Table 1), a tendency for disease onset to be delayed with increasing age at immunisation in the dystrophic rats was found. Animals sensitised at 60 days of age developed EAU in black-eyed RCS dystrophic and non-dystrophic rats. The figure displays the relationship between the proportion of affected eyes, and time after sensitisation with 50 µg bovine S-antigen in FCA together with B pertussis inoculation (see Methods).

<table>
<thead>
<tr>
<th>Age at sensitisation (days)</th>
<th>Dystrophic rats</th>
<th>Non-dystrophic rats</th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>18/9 (9/9)</td>
<td>26/1 (5/5)</td>
</tr>
<tr>
<td>85-90</td>
<td>22/3 (8/3)</td>
<td>26/0 (10/5)</td>
</tr>
<tr>
<td>105</td>
<td>30/5 (8/5)</td>
<td>ND</td>
</tr>
</tbody>
</table>

* n=90 no of eyes

Onset of EAU in dystrophic rats sensitised at 105 days was delayed in comparison with rats sensitised at 60 days (p=0.003, Wilcoxon rank sum test).
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Table 2  Clinical features of uveoretinitis in dystrophic and control strain RCS rats

<table>
<thead>
<tr>
<th>Stage of disease</th>
<th>Prevalence of clinical features (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0-5 days early</td>
</tr>
<tr>
<td>Retinal detachment</td>
<td>D  C  D  C  D  C</td>
</tr>
<tr>
<td>Anterior uveitis</td>
<td>52  3  0  1  0  6</td>
</tr>
<tr>
<td>Retinal vasculitis</td>
<td>0  26  0  12  0  0</td>
</tr>
<tr>
<td>Pigmentary changes</td>
<td>44  38  61  68  35  71</td>
</tr>
</tbody>
</table>

D = dystrophic strain; C = control strains.
Fifteen animals of each strain were examined on three different days during the three time intervals. The prevalence of individual features of disease is expressed as a percentage of 90 observations at these three stages.

uvectis with a mean onset of 19 days, while those sensitised at 105 days of age developed disease at a mean time of 31 days (p=0.003, Wilcoxon rank sum test). No such tendency was seen in the control animals. It was concluded that whereas both dystrophic and non-dystrophic rats were susceptible to disease, the dystrophic strain showed a delay in developing disease with increasing age at sensitisation.

CLINICAL FEATURES OF DISEASE IN DYSTROPHIC VS NON-DYSTROPHIC CONTROLS

All animals showed retinal inflammation by indirect ophthalmoscopy; patchy retinal infiltrates, venous dilatation, tortuosity, periphlebitis, and disc oedema. Disc oedema persisted for a few weeks after other signs of retinal vasculitis has subsided. The fundal view was not clouded by vitritis, but severe anterior uveitis prevented fundoscopy for a few days in some animals. The fundal appearances were similar in the two strains and resembled the previously reported changes in the Lister rat. Abnormalities of retinal pigmentation, consisting of patchy hyperpigmentation and atrophy, developed in both dystrophic animals and non-dystrophic controls. There was no difference in the age of onset of pigmentary changes between animals of the two strains, whether sensitised or not (mean age at onset 123 days, 95% confidence interval 117–129).

There was a striking difference in the initial phases of disease between dystrophic and non-dystrophic controls (Table 2): 17/30 eyes of the control strain developed anterior uveitis with miosis and synchiae formation as the initial presentation of the uveitis. In contrast, anterior uveitis was not seen in any of the dystrophic animals (χ² = 23.7, P<0.001).

Early disease in the dystrophic strain was characterised by the sudden onset of serous retinal detachment which subsided after a few days; this was followed by other signs of retinal vasculitis similar to those seen in the control strain. Eighty per cent (24/30) of dystrophic eyes developed total serous retinal detachments in the first week of disease compared with 10% (3/30) of control strain eyes (χ² = 29.7, P<0.001).

FLUORESCIN ANGIOGRAPHY AND HISTOLOGY

Fluorescein angiograms showed leakage from the optic disc and areas of intraretinal infiltrates during active disease: no such leakage was seen prior to the development of clinical signs (Fig 3). There was no difference in the angiographic appearances between dystrophic and control strains, and appearances were again similar to those previously reported in the Lister rat. During the phase of serous retinal detachment there was widespread, diffuse leakage of fluorescein in the subretinal space.

Unsensitised dystrophic rats showed the reported changes of photoreceptor dystrophy with extensive loss of the photoreceptor cells and the spontaneous accumulation of debris in the subretinal space (Fig 4A).

All sensitised animals developed an extensive mononuclear infiltrate surrounding the retinal vessels, but minimal choroidal inflammation was seen. The optic disc region showed gross oedema and leukocytic infiltration. In non-dystrophic animals that had developed mild clinical disease, there were focal retinal infiltrates but the surrounding retinal architecture remained normal.

Figure 3  Fundus photographs and fluorescein angiograms of dystrophic rats showing clinical features of EAU. (A) serous retinal detachment; (B) normal optic disc; (C) swollen optic disc with dilated retinal vessels; (D) peripheral retinal phlebitis; (E) and (F) are angiograms corresponding to the areas shown in (C) and (D), demonstrating dilated vessels and capillary and venous leakage.
In the dystrophic animals leucocytic infiltration into the retina was characteristically diffuse rather than focal and of greater severity than in the non-dystrophic strain. In the late stages of EAU dystrophic animals showed loss of subretinal debris with thinning of the neural retina; a small inflammatory infiltrate remaining in the inner retina at a time when funduscopic evidence of disease was no longer evident (Fig 4C). In late EAU there were extensive areas of gliosis accompanied by neovascularisation in the dystrophic retina (Fig 4D). Although similar changes could be seen in the control strain, they were much less widespread than in the dystrophic strain. In the anterior segment there was mononuclear cell infiltration of the ciliary body and occasionally evidence of posterior synechiae. In an earlier series of experiments, animals were sacrificed during the early stages of disease. Figure 4E shows the detached retina of a dystrophic rat sensitised according to an identical protocol at the age of 60 days. The animal was sacrificed 14 days after sensitisation and 3 days after the onset of extensive serous detachments. Large clumps of granulocytes and mononuclear cells are seen in the subretinal space.

Discussion
In this study we have demonstrated that the RCS rat is susceptible to EAU after sensitisation with retinal S-antigen, though older animals with photoreceptor dystrophy show a delay in disease onset. The presence of photoreceptor dystrophy was associated with a marked alteration in the form of early clinical disease from an acute anterior uveitis to that of a complete serous retinal detachment. As the two strains of rat differed only by the presence or absence of the photoreceptor dystrophy gene(s) we presume that these differences in disease expression were not due to differences in immune responsiveness or to differences in ocular vascular anatomy between the strains. In general the photoreceptor layer is the site of the most profound T lymphocyte infiltrate and retinal cell destruction in EAU, while in the Lewis rat initial T lymphocyte infiltration occurs in the ciliary body. The route and severity of early cellular infiltration into the eye in EAU varies with the ocular vascular anatomy, the dose of antigen, the composition of adjuvant as well as between strains of the same species. In EAU, fluorescein angiography and electroretinography demonstrate that the cellular infiltrate is preceded by a period of increased retinal vascular permeability and abnormal photoreceptor function, and there is
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Figure 4 Histology of posterior segment changes in EAU produced in dystrophic and non-dystrophic RCS rats (haematoxylin and eosin). (A) Section from non-sensitised dystrophic rat, 60 days old, showing extensive loss of photoreceptor cells and their replacement with amorphous debris (×130). (B) Section from non-dystrophic rat 86 days after sensitisation. There is a focal outer retinal infiltrate with adjacent disruption of the RPE. The surrounding retina shows normal photoreceptor layer morphology (×150). (C) Section from dystrophic rat with clinically inactive disease, 86 days after sensitisation. There is total loss of photoreceptor layer and associated debris. The inner retina is disrupted and there is irregularity of pigmentation at the RPE (×150). (D) Area of neovascularisation in dystrophic rat, 86 days after sensitisation (×150). (E) Section taken during the phase of acute retinal detachment, 14 days post-sensitisation in a dystrophic rat. There is a marked serous elevation of the retina with subretinal aggregates of mononuclear cells and granulocytes. A mild infiltrate is present in the outer retina (×66).

some evidence that humoral factors may be responsible for those early changes.12–19 Retinal detachment is known to occur during EAU in species other than the rat19 where it is usually adjacent to areas of choroiditis and to RPE proliferation; and where the subretinal fluid contains lymphocytes and macrophages derived from the choriocapillaris. Extensive retinal detachment with subretinal granulocyte exudation can occur in hyperacute EAU in the Lewis rat, induced by high-dose sensitisation with S-antigen, where it is considered to be immune-complex mediated.16

The retina of the RCS rat has three characteristics which may predispose it to develop retinal detachment so readily in EAU. Firstly, normal adhesion of the retina to the pigment epithelium is compromised by the loss of interdigitations between the photoreceptors and RPE, as the outer segments degenerate and
debris accumulates between the photoreceptor layer and the RPE. Secondly, the interphotoreceptor matrix, which promotes retinal adhesion, is abnormal early in the course of the dystrophy.20 Thirdly, both the outer and inner blood-retinal barriers become spontaneously leaky to plasma protein as early as 21 days of age.21 In EAU these characteristics may allow the early accumulation of large amounts of subretinal fluid and subsequent detachment prior to the maximal influx of inflammatory cells, so enabling sufficient antibody to enter the subretinal space to produce a local Arthus reaction.

We did not find evidence of altered susceptibility to EAU with increasing age at sensitisation in the dystrophic rats though there was a tendency for EAU onset to be delayed in the older animals. There is known to be a reduced susceptibility to, and reduced severity of EAU in animals when photoreceptor outer segment turnover is reduced by disturbance of the normal day/night cycle. Thus, rearing in constant light (which induces photoreceptor degeneration) or constant dark22 (which moves S-antigen to the inner portions of the photoreceptor layer) will reduce the severity of EAU as well as reduce the amount of S-antigen in the photoreceptor outer segments. We find that constant dark rearing also delays the onset of EAU even in the high responder Lewis strain of rat (E Kasp and S Suleyman, unpublished data). These findings contrast with the exacerbation of disease severity brought on by constant dark, in the collagen induced arthritis model of autoimmune disease.23

In the present experiments, where RCS photoreceptor metabolism is disturbed because of the RPE abnormality, it is difficult to know whether the amount of S-antigen available to sensitised recirculating T-lymphocytes is reduced because of the reduced number of photoreceptors, or increased because of the disruption to the blood-retinal barrier. The tendency for an earlier onset, and shorter course of disease in the dystrophic strain, would be expected if there was a relatively greater access to a reduced amount of the target antigen.

Altered expression of EAU in the dystrophic strain might also arise from innate sensitisation to photoreceptor proteins, known to occur in both animals and humans with photoreceptor dystrophies. The RCS rat spontaneously develops IgM antibodies against photoreceptors in the early stages of dystrophy24 and a small minority develop raised levels of antibody against S-antigen itself.25 Raised levels of antibodies to photoreceptors have been detected in humans with retinitis pigmentosa (RP)26 and the frequency of retinal-specific B cells (found after Epstein-Barr virus transformation) is higher than normal.27 In RP patients T suppressor cells are reduced and there are slight increases in interleukin-2 receptor expression or peripheral lymphocytes and in HLA class II expression on RPE cells.28 These changes, taken together, suggest that in the RCS rat as well as in retinitis pigmentosa the immune system is primed for autoimmune reactions against the retina, which may well contribute to the disease process. Antiretinal antibodies are also known to be associated with increased retinal vascular permeability in a variety of degenerative and inflammatory retinal disorders in man.29

In conclusion, this model of EAU demonstrates that a genetically determined biochemical dysfunction of target photoreceptor cells influences the clinical expression of retinal disease produced by peripheral autosensitisation to S-antigen. Further characterisation of the vascular and cellular events in early EAU in the RCS dystrophic rat may help to explain how these genetic factors could be operating at the molecular level.

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