Age-Related Macular Disease

Age-Related Macular Disease: Aetiology and Clinical Management

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Caucasian Communities

Age-related macular disease (AMD) accounts for about 50% of registered blindness in England and Wales, and this high prevalence is likely to exist in all economically developed Caucasian communities. A recent analysis indicates that the prevalence appears to be increasing at a rate not fully explained by the increasing age of the population, and, as a cause of visual loss, AMD is as common now as diabetes and glaucoma during working life. Despite the early expectations, it is now evident that the present techniques of laser treatment will not have a major impact on blindness due to AMD. Other forms of treatment are under trial, such as ionising radiation and photodynamic therapy, but they may not be vastly more successful than photocoagulation. This means that our knowledge of the behaviour and pathogenesis of the disorder must be re-examined in the hope that alternative approaches to management can be identified.

Twin and sibling studies provide good evidence of genetic predisposition in AMD, and it is believed that the predisposition becomes manifest in the presence of appropriate environmental influences. It is likely that more than one gene is involved, although the number is probably small. In these respects AMD is similar to other complex traits. It follows that ageing at the macula would vary qualitatively within a community, occurring in those with the most severe genetic predisposition and environmental pressures. The high prevalence of clinically detectable age-related maculopathy (ARM) in those over the age of 65 years implies that the causative genes are common in industrial societies.

Non-Caucasian Communities

In the past AMD was rarely recognised in Japan. However, disability due to macular disease in middle or late life, and hospital referrals with this disorder, suggest that AMD has become common in the last two decades, at least in urban communities. A similar trend is appearing in other parts of Eastern Asia. There is also a strikingly high prevalence of macular disease in elderly Inuit in Greenland.

The phenotype (clinical appearance) of AMD appears to vary in different communities. In Japan the impression is that polyoidal choroidopathy is common, whereas, in Caucasians, the growth of new vessels is the most prevalent complication. In the Inuit population the characteristic process is atrophy. In both Inuit and Japanese the visual loss occurs without there being pre-existing recognisable age changes at the level of Bruch’s membrane, such as soft drusen. In neither has genetic predisposition been sought so far, although the increasing incidence implies environmental factors are important.

Future Research

Cross sectional studies in populations with different genetic backgrounds but similar environment, and populations with similar genetic background with different environments, would serve to prove that both influences are important.

Future research seeking the abnormal genes is likely to be rewarding. The techniques of linkage disequilibrium and sibling pair analysis have been well worked out, and have met with success in other complex disorders such as multiple sclerosis, diabetes, bipolar illness and schizophrenia. Both investigative techniques benefit from knowledge of candidate genes. Candidate genes may be found by recognition of associated characteristics, such as loss of iris colour, or identification of genes causing disorders with phenotype similarities to AMD such as Sorsby fundus dystrophy and Doyne macular dystrophy, together with knowledge of the disease processes.

In Caucasians visual loss results from choroidal neovascularization, detachment of the retinal pigment epithelium (RPE) or geographic atrophy. It is widely believed that these occur in response to accumulation of debris in Bruch’s membrane, which is recognised clinically as drusen and pigmentary changes referred to as age-related maculopathy (ARM). It is believed that the debris in Bruch’s membrane is derived from the RPE, which discharges cytoplasmic material throughout life into the inner portion of Bruch’s membrane in order to achieve cytoplasmic renewal. It is likely that the material is cleared through the choriocapillaris. Some information exists concerning age changes in the RPE, Bruch’s membrane and choroid from laboratory studies of donor eyes. However, the inter-relationship between age change in the different tissues is not established. Clinical and laboratory studies imply that the quantity, distribution and chemical composition of the debris determine both the magnitude of risk, and the type of lesion causing visual loss. From these observations it is possible to speculate on the potential genetic influences that may modulate ageing. Increased outer segment turnover may explain the high levels of RPE autofluorescence in diseases due to mutations in the RDS gene. A similar effect may occur if there is reduced activity of RPE degradative enzymes, or free radical damage to the substrate of degradation. Age change in Bruch’s membrane such as cross linkage of collagen would predictably accelerate accumulation of debris. The mechanisms whereby material is cleared from Bruch’s membrane may also be under genetic influence. Considerable variation of age changes exists from one donor to another of a similar age, reflecting the complexity of genetic and environmental influences.

Thus, there are potentially many candidate genes influencing ageing at the macula. Whether or not a single gene determines risk in an individual or family is unknown, although the reputed high gene frequency in the population implies that a single gene should not be assumed.

Of the environmental influences smoking has most consistently been associated with increased risk, but surprisingly not all studies have demonstrated this. Dietary intake of carotenoids, other micronutrients and hypertension have also been implicated. However, none of these readily explain the apparent increase in prevalence of
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Introduction

Age-related maculopathy (ARM) is a degenerative disorder of the central retina typically with an age of onset after the fifth decade. It is characterised in the early stages by drusen, pigmentary changes and degeneration of the retinal pigment epithelium (RPE). In the later stages there is atrophy of the photoreceptors and RPE (geographic atrophy or dry form) and choroidal neovascularisation (CNV or wet form); the latter resulting in the typical disciform scar. Only the later stages of ARM (termed AMD) featuring geographic atrophy and/or CNV result in moderate or severe loss of vision.

Some of the hypotheses for the development of CNV are that
(a) tissue barriers to blood vessel growth are disrupted by degeneration
(b) inflammatory cells incite the neovascular response
(c) decreased choroidal perfusion and impaired oxygen transport result in ischaemia and neovascularisation

The only proven treatment for choroidal neovascularisation is focal laser photocoagulation. Less than 20% of lesions are eligible for laser treatment at clinical presentation and even in these the benefit is modest. Thus, many different therapeutic approaches are being investigated which include preventive strategies and methods to manage established disease.

Preventive Strategies

CNV in the expanding elderly population brings a high level of visual disability and social cost. Thus, there is great interest in preventing the ingrowth of new vessels before there is severe loss of visual function, particularly since there is no treatment for the atrophic form and existing therapies are of minimal benefit for CNV.

A number of preventive strategies are therefore under consideration. As oxidative stress has been suggested as playing a role in macular tissue damage leading to ARM, dietary supplementation with antioxidants is being tested in a randomized controlled trial. The National Eye Institute, USA, is sponsoring the Age-Related Eye Disease Study (AREDS); a multicentre, clinical trial to evaluate the role of antioxidant vitamins (with or without zinc supplementation) in the prevention of age-related macular degeneration and cataract. As the development of these degenerative disorders is often a long process, any beneficial effects in the treated group are unlikely to become apparent for at least another 5 years.

Within the past several years, there has also been significant interest in the role of low intensity laser treatment to eyes with large diffuse drusen in the prevention of CNV and loss of vision. Pilot studies have reported a decreased incidence of CNV and a lower rate of loss of vision among treated eyes. Initial results from a large randomized clinical trial (The Choroidal Neovascularization Prevention Trial, CNVPT) are now available. Essentially this study enrolled patients into one of two groups. The first group consists of patients with established neovascularisation in one eye with soft drusen in the fellow eye (Fellow Eye Study). The second group consists of individuals with bilateral soft drusen (Bilateral Drusen Study). Photodynamic active argon laser photocoagulation was performed in the fellow eye in the Fellow Eye Study and in one eye in the Bilateral Drusen Study. Interim analysis has demonstrated increased rates of CNV but decreased rates of vision loss in fellow eyes treated with laser in the Fellow Eye Study. Among patients with bilateral drusen, treated eyes and observed eyes had similar rates of CNV and loss of vision.

Thus, there is a possibility that some interventions may succeed in reducing the rate of visual loss but all the indicators suggest that the effects, if any, will be modest.

Current Status of Clinical Trials for CNV of AMD

While prevention remains a primary goal, a significant proportion of the ageing population exhibits fully developed ARM. Various new therapeutic approaches are under investigation in their management. These include the use of photodynamic...