The cardiototoxicity of eosinophils

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Summary

Although an association between high blood eosinophil counts and endomyocardial disease has been known for nearly a hundred years, the reasons for this were not understood. Brockington, Luzzatto and Osunkoya (1970) suggested that eosinophil leucocytes, in susceptible persons, by some unknown mechanisms, cause endomyocardial damage. Evidence to support this possibility has come from three sources: (1) Clinical studies have shown that very high blood eosinophil counts, from any cause, can be associated with endomyocardial disease, and in some patients it has been possible to show that eosinophilia preceded the onset of heart disease. (2) The development of heart disease has been associated with the presence of degranulated eosinophils in the blood and tissues, including damaged endomyocardium, and raised serum levels of eosinophil granule basic proteins have been found in many of these patients. (3) Low concentrations of eosinophil secretion products (which contain these eosinophil granule basic proteins) have been found to injure isolated heart cells in vitro. Studies with purified eosinophil granule basic proteins have shown that cardiac cell damage is the result of a specific toxic effect of eosinophil cationic protein on the plasma membrane and two enzyme complexes (pyruvate dehydrogenase and 2-oxoglutarate dehydrogenase) involved in mitochondrial respiration. These results support the suggestion that under certain conditions, eosinophils may damage the heart, leading to endomyocardial disease, and they offer new approaches for the early diagnosis and treatment of endomyocardial disease both in temperate and tropical countries.

KEY WORDS: endomyocardial disease, eosinophilia.

Introduction

There have been clinical reports of an association between high blood eosinophil counts and endomyocardial fibrosis for nearly a hundred years, but wide recognition of this was only achieved following the report of two patients studied during life by Löffler (1936). Brockington et al. (1970) described a patient with eosinophilic leukaemia and endomyocardial fibrosis, and were so struck by the association that they suggested that eosinophil leucocytes, in susceptible persons, by some unknown mechanisms, cause endomyocardial damage. Zucker-Franklin (1971) also raised the possibility that eosinophils might damage the heart by analogy with carcinoid-induced endocardial lesions. Independently, Yam et al. (1972) noted the frequent association of endocarditis and eosinophilic leukaemia, and felt that this could not be coincidental. They also proposed that there was a cause-effect relationship similar to that of cardiopulmonary lesions in metastatic carcinoid syndrome. In addition, they pointed out that endomyocardial fibrosis had occurred in patients who had a prolonged eosinophilia from many different causes. The possibility that eosinophils might release their toxic constituents and injure the endocardium did not receive wide acceptance, largely because it was held at that time that eosinophils had a modulating or damping role in inflammation. The later demonstration that eosinophils could kill some parasites in vitro initiated a large amount of subsequent work, which has shown that eosinophils have a marked capacity to injure some organisms and cells (Spry, 1978).

The purpose of this paper is to review recent clinical and experimental studies which have shown that eosinophils have a potent cardiototoxic capacity, which may explain the well-known association between high blood eosinophil counts and endomyocardial fibrosis. Much of this work has been done with purified eosinophils (and their constituents) from patients with the hypereosinophilic syndrome (Spry, 1982; Spry et al., 1983a), over 80% of whom developed eosinophilic endomyocardial disease (Parillo et al., 1979; Davies et al., 1983b). It has been
suggested that a similar pathogenetic mechanism is responsible for the development of tropical endomyocardial fibrosis (Olsen and Spry, 1979), based on the observation that eosinophilic and tropical endomyocardial fibrosis are pathologically indistinguishable (Brockington and Olsen, 1973). If this is so, then there are important implications for therapy and prevention of endomyocardial disease in many patients both in temperate and tropical climates.

Patients and methods

Clinical features of the patients have been described elsewhere (Spry et al., 1982; Davies et al., 1982). In brief, there were 15 patients with the hypereosinophilic syndrome, 11 of whom had biventricular eosinophilic endomyocardial disease. Five patients presented with heart disease, and 6 others developed cardiac involvement, a mean of 1-3 years after hypereosinophilia was first detected. Nine had mitral valve disease and 6 had tricuspid regurgitation. Cardiac histology showed acute necrotic lesions in 4 patients and late fibrotic lesions in 5. The mean blood eosinophil count at presentation was 20·1 x 10⁹ /litre. A mean of 3·7 x 10⁹/litre degranulated blood eosinophils (Spry and Tai, 1976) were found in the blood of 10 patients.

Measurements were made of the capacity of patients' blood eosinophils to injure isolated rat heart cells in vitro, as described previously (Tai et al., 1982). Rat heart cells were isolated by collagenase and hyaluronidase digestion, and incubated with supernatants from purified eosinophils which had been stimulated with zymosan-C₄b. Purified eosinophil and neutrophil granule proteins were kindly provided by Dr Inge Olsson, Lund, Sweden (see Venge and Olsson, 1980). The O₂ uptake of isolated rat heart cells was measured with an O₂ electrode, and the integrity of the mitochondrial respiratory chain was assessed using isolated rat heart cell mitochondria (Tai et al., 1982).

The concentration of eosinophil granule basic proteins in serum was assayed using a mouse monoclonal antibody (EG2) which detected a common epitope on eosinophil cationic protein (ECP) and eosinophil neurotoxin (EPX). These two proteins were assayed together in serum, using this monoclonal antibody, in a competitive inhibition enzyme immunoassay with peroxidase-linked rabbit antimouse immunoglobulin (Tai and Spry, unpublished).

Results

Blood eosinophil counts

All 11 patients with eosinophilic endomyocardial disease presented with hypereosinophilia (blood eosinophil counts more than 1·5 x 10⁹/litre), but 3 patients had normal blood eosinophil counts for several weeks or months during their illness. In one patient, the eosinophil count became normal spontaneously. In 2 other patients, normal counts developed after steroid treatment had begun, and in one of them, the eosinophil count has remained normal for 18 months, except for short periods when it rose above 1 x 10⁹/litre.

Degranulated blood eosinophils

Twelve of the 15 patients with the hypereosinophilic syndrome had more than 0·5 x 10⁹/litre degranulated blood eosinophils. Eleven of these patients were shown to have eosinophilic endomyocardial disease. One patient with endomyocardial disease had only intermittently raised blood eosinophil counts and few degranulated blood eosinophils.

Another patient, who had many degranulated blood eosinophils at the start of his illness, has subsequently been found to have a normal heart, judged clinically and by echocardiography, and degranulated cells have disappeared from his circulation. No degranulated blood eosinophils have been seen in blood films from the 2 other patients with hypereosinophilia without endomyocardial disease.

One patient showed a remarkable association between a rise in the number of degranulated blood eosinophils and the development of acute endomyocardial disease. He had had blood eosinophil counts of between 5 and 10 x 10⁹/litre for 2·5 years with no evidence of endomyocardial disease. Then he became ill with fever, exhaustion, cough, abdominal pain and rapid weight loss. The eosinophil count rose to more than 100 x 10⁹/litre, and for the first time, many of them were found to be degranulated. The heart enlarged and cardiological investigations now showed that he had developed bi-ventricular endomyocardial disease in the acute necrotic stage. He was treated with steroids which produced a rapid improvement in systemic and cardiological symptoms and signs. Several months later he was able to return to work, and echocardiography and endomyocardial biopsy have shown that his disease has not progressed from the acute necrotic stage.

The mean concentrations of serum ECP and EPX together, in 13 patients with hypereosinophilic syndrome without heart disease was 2030 ± 2900 ng/ml (mean ± s.d.). The mean level in 9 patients with eosinophilic endomyocardial disease was significantly higher: 4540 ± 2600 ng/ml.

Injury to rat heart cells in vitro

Supernatants from eosinophils which had been stimulated with C3b gave rise to both a dose-dependent cytotoxic effect and a stimulation of O₂
uptake by isolated rat heart cells in vitro. This effect on respiration was blocked by ouabain, showing that it occurred as a result of an increase in the activity of Na⁺/K⁺ ATPase in the plasma membrane. These eosinophil supernatants also inhibited pyruvate dehydrogenase and 2-oxoglutarate dehydrogenase in isolated rat heart mitochondria. Neutrophil secretion products did not have these effects. Preliminary studies with purified ECP, and a mouse monoclonal antibody which binds to ECP and EPX, have suggested that ECP was responsible for the cytotoxicity in eosinophil supernatants. Single cell preparations from other tissues were not injured by eosinophil supernatants, and equal concentration of other unrelated proteins, with a similar basic charge, had no effect on heart cell viability or respiration or isolated mitochondrial enzymes (Tai et al., 1982).

Discussion

This study was designed to review the evidence that eosinophils could damage the heart and give rise to endomyocardial disease. Three separate lines of evidence support this view. The first is derived from clinical studies on patients with hypereosinophilia. The second is related to the detection of degranulated eosinophils in the blood and tissues, and the third is the direct demonstration that eosinophil granule proteins can injure isolated heart cells in vitro.

Clinical studies

These have provided some of the most compelling evidence that eosinophils may cause endomyocardial disease. Two reports have listed the wide range of clinical disorders in which hypereosinophilia may be associated with endomyocardial disease (Olsen and Spry, 1979; Schooley et al., 1980). The most likely common factor in these diseases was the presence of a marked blood eosinophilia which, in many patients, clearly preceded the onset of clinical evidence of heart disease. The temporal relationship between eosinophilia and heart disease was clearly shown in 6 of the 11 patients with endomyocardial disease reported here. It is interesting to consider why some patients present both with heart disease and hypereosinophilia, and why others only develop heart disease later. The most likely explanation is that this type of heart disease is asymptomatic in its early stages unless there is a widespread and severe carditis. For this reason most patients will only present with early heart involvement if they have other complications of the underlying disease which is producing the eosinophilia. This may explain why it has been impossible as yet to detect and define the early stages of tropical endomyocardial fibrosis. Patients who develop a moderate eosinophilia in areas of endemic parasitic disease, including filariasis, may not be distinguishable from other parasitized individuals until their heart disease is in a late stage, when blood eosinophil counts may have returned to normal. The disappearance of high blood eosinophil counts has been documented in patients with eosinophilic endomyocardial disease, and these patients appear to be an intermediate group between patients with the tropical and eosinophilic forms of endomyocardial disease.

Degranulated blood eosinophils

The presence of degranulated blood eosinophils in patients with hypereosinophilia and endocardial thrombi and fibrosis has been known for over 60 years (Shapiro, 1919). It was suggested (Spry and Tai, 1976) that these cells had released their granule contents which damage the endomyocardium leading to the development of endomyocardial disease. This hypothesis has been supported by studies on the patients reported here. Ten patients with eosinophilic endomyocardial disease had many degranulated blood eosinophils in their circulation. Unfortunately, in one patient with acute necrotic cardiac lesions, eosinophils were not available for study. In one other patient, although degranulated cells were found for 3 months, there was no clinical evidence of endomyocardial disease. As degranulated cells then disappeared from the blood, invasive cardiac studies (which might have shown early lesions) were not carried out. The clinical significance of the presence of degranulated eosinophils in blood smears was confirmed by Jaski et al., (1978) who noted the presence of these cells during life in a patient with carcinoma of the lung-induced hypereosinophilia, who was shown to have endomyocardial lesions at post-mortem. We have studied a similar patient, and documented an identical sequence of events.

The recent demonstration of degranulated eosinophils in both the eosinophilic and tropical forms of endomyocardial disease (Olsen, 1983) raises the possibility that degranulation may occur locally in the heart, and degranulated eosinophils may not be present in the circulation of these patients. This important finding strengthens the view that both forms of the disease have a common pathogenesis related to the presence of eosinophil granule basic proteins in the heart. This view will be strengthened if it can be shown that these toxic proteins are present in endomyocardial biopsies from patients in early stages of both forms of the disease.

Raised levels of eosinophil granule basic proteins were found in the circulation of patients who had many degranulated blood eosinophils. The significance of this finding is not clear as patients with an eosinophilia (which was the result of parasitic diseases) and who did not have endomyocardial disease
have been found to have equally high blood eosinophil granule basic protein levels (Tai, unpublished). It is possible that circulating eosinophil granule basic proteins have an initial toxic effect on the heart, causing eosinophils to localize there and degranulate. Alternatively, a small proportion of patients who develop hypereosinophilia may lack effective inhibitors for the potentially cardiotoxic effects of eosinophil granule basic proteins. This latter explanation could account for the low incidence of endomyocardial disease in patients with asthma-induced hypereosinophilia (Olsen and Spry, 1979).

Heart cell injury in vitro

Human eosinophil granule basic proteins have been shown to injure rat heart cells in vitro, and inhibit isolated mitochondrial respiration. Less than $10^{-9}$ mol/litre eosinophil granule basic proteins produced these effects, and these concentrations are likely to be achieved within the heart in vivo. Due to their strong charge, eosinophil granule basic proteins are likely to bind closely to cell membranes. This may be the first step in membrane damage which alters permeability to sodium. It is not known whether these basic proteins enter cardiac cells, but if they do so, they are capable of inhibiting mitochondrial respiration directly. Either (or both) of these events are likely to lead to cardiac cell death.

Previous studies, which attempted to show a toxic effect of eosinophils on cardiac cells, met with little success (Epps and Bankhurst, 1978; Parrillo and Fauci, 1978). $^{51}$Cr labelled target cells, including cultured atrial cells, were treated with antibody and/or C3b and incubated with blood eosinophils. Only a small proportion of the chromium was released. However, neither study measured the concentration of eosinophil granule proteins which were released in these experiments, and this may have been small. The results do, however, suggest that antiheart antibody, or direct complement activation on the surface of cardiac cells, are unlikely to be involved in the induction of eosinophil dependent endomyocardial damage. This is supported by studies on patients' sera which have failed to detect antiheart antibody (Davies, unpublished). However, it should be noted that blood eosinophils which have little initial cytotoxic capacity in normal people may be activated in vitro, and eosinophils in patients with an eosinophilia are metabolically and functionally more effective than normal. These changes are linked to alterations in oxygen metabolism (Pincus et al., 1981), and it is possible that the production of oxygen radicals augments the cytotoxic effects of the eosinophil granule basic proteins (Olsen and Spry. 1979).

Two important questions remain to be answered: Why is the endocardium so susceptible to eosinophil-dependent injury, and what determines the rate at which endocardial disease progresses to the late fibrotic stage? Selective injury to the endocardium may be related to regional differences in cardiac blood supply, metabolism or capacity to repair injury. Alternatively, toxic eosinophil granule basic proteins may be concentrated in the endocardium. Each of these possibilities can now be studied, as a model of eosinophil granule basic protein dependent cardiac injury and methods to do so are being developed in rats (Davies et al., unpublished).

The second question is more difficult to approach, but it is of considerable importance as it is now likely that tropical endomyocardial disease is a similar, but more slowly progressing, form of eosinophilic endomyocardial disease (Davies et al., 1983b). The late fibrotic lesions in both forms of the disease are identical (Brockington and Olsen, 1973), and it has been thought for many years that they might have a common pathogenesis related, in some way, to the presence of eosinophils in blood or tissues (Olsen and Spry, 1979). Unfortunately, it is impossible to study mechanisms of chronic injury using isolated heart cells in vitro, but it may also be possible to approach this question using the rat model of eosinophil-dependent injury. Although these issues, and other questions, such as the geographical distribution of tropical endomyocardial fibrosis, remain to be resolved, studies on the interaction between eosinophils and heart muscle have provided many new approaches to the study of both eosinophilic and tropical endomyocardial disease. It is hoped that this will provide further insights into the mechanisms of this disease and will lead to effective methods for prevention, early diagnosis and treatment.

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References


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