Extralymphatic disease due to bancroftian filariasis

G. Dreyer¹,² P. Dreyer² and W.F. Piessens³

Abstract

Infection with Wuchereria bancrofti, Brugia malayi, or B. timori not only affects the structure and function of lymphatic vessels but is also associated with extralymphatic pathology and disease. Because it is now possible to detect living adult worms by ultrasonography, much emphasis is placed on lymphatic pathology. However, the finding of renal damage in asymptomatic microfilaremic carriers has led to increased recognition of the importance of extralymphatic clinical manifestation in bancroftian filariasis. The authors present a number of clinical syndromes that may be manifestations of extralymphatic filarial disease and discuss possible mechanisms that cause these conditions. The main purpose of this paper is to raise the awareness of students and physicians of the prevalence and the importance of extralymphatic disease in bancroftian filariasis so that it is diagnosed and treated properly and also to alert for the need of additional research in this area.

Introduction

Lymphatic filariasis persists as a major cause of clinical morbidity and a significant impediment to socioeconomic development in Asia, Africa and the Western Pacific, as well as in certain regions of the Americas. It is estimated that at least 120 million persons are infected with W. bancrofti and B. malayi. Of these, 40 million patients suffer from lymphedema, elephantiasis or hydroceles - the well-known clinical manifestations of lymphatic disease (1).

Recently, dramatic advances in our understanding of the pathogenesis of this disease have led to the recognition that subclinical lymphatic damage is present in all infected patients. Lymphatic dysfunction resulting from this damage predisposes to local microbial infection, which in turn exacerbates lymphatic pathology (2-5). However, little attention has been paid in recent years to clinical manifestations of extralymphatic disease caused by filariasis. This paper summarizes the current knowledge about this topic. It is intended to reemphasize the importance of extralymphatic morbidity in bancroftian filariasis, and to highlight the gaps in our knowledge of the pathogenic mechanisms that underlie the various clinical syndromes.

The extralymphatic syndromes resemble clinical entities of nonfilarial origin, and it is often impossible to establish with absolute
certainty the filarial etiology of extralymphatic disease manifestations in an infected individual. Similar diagnostic uncertainties also apply to some of the “classical” manifestations of lymphatic filariasis (4,5). Unlike lymphatic disease syndromes, the extralymphatic manifestations of bancroftian filariasis are not caused by adult worms per se, but by microfilariae, by diffusible products from as yet undefined parasite stages, or by immune complexes. Extralymphatic filarial disease is thus heterogeneous in its pathogenesis and clinical manifestations.

**Arthritis**

Arthritis has long been recognized as a possible manifestation of filarial infections (6). Two clinical types have been described. The first is oligoarticular “filarial arthritis”, a rare condition that typically affects just one large joint, most commonly a knee. The entity can be distinguished from other forms of arthritis by clinical criteria associated with laboratory findings and by its prompt response to diethylcarbamazine (DEC) (7). Its pathogenesis is unclear. Synovial fluid from the affected joint ordinarily does not contain microfilariae, adult worms or pyogenic organisms, but the monoarticular inflammation may reflect a tissue reaction to a filarial worm in the vicinity of the joint. Rarely, lymphatic fistulation into the synovial sac causes chylous arthritis (8).

The second manifestation, polyarticular “filarial pseudo-rheumatism”, appears to be less common in lymphatic filariasis than in onchocerciasis, loiasis or mansonielsis. Its pathogenesis is believed to involve intra-articular deposition of immune complexes (9). However, immune complexes containing filarial antigens have yet to be detected in synovial tissue, and they are present in sera from many infected patients without arthritis (10,11). Intact microfilariae have been detected intra-articularly in some patients with filarial polyarthritis and in animals (12). In such cases, local release of proteases by the worms (13) may directly damage synovial tissue.

**Renal disease**

Despite case reports over the years describing instances of glomerulonephritis, hematuria and proteinuria in patients with filarial infections, the prevalence of renal abnormalities in bancroftian filariasis has only recently been formally determined. Renal disease is rarely observed in patients without microfilaremia. In contrast, ~45% of untreated microfilaremic patients have renal pathology manifested as microscopic hematuria (~35%) and/or proteinuria (~20%). Treatment with DEC exacerbates the proteinuria in these patients and also triggers transient hematuria or proteinuria in many microfilaremics persons with a normal urine analysis prior to treatment (14). Mechanical damage to glomeruli by circulating microfilariae is believed to account for hematuria in some cases (15,16), but damage caused by deposition of immune complexes in the glomerular basement membrane is likely to be a far more common cause of renal pathology in bancroftian filariasis. Such complexes have been observed in renal biopsies of patients with filarial infections and glomerulonephritis (17–19). However, only in patients with onchocerciasis has it been formally determined that antigens within immune complexes deposited in the kidneys are of filarial origin (20).

**Tropical pulmonary eosinophilia**

Several helminth species can cause pulmonary infiltrates and eosinophilia (PIE), a transient syndrome resulting from the migration of worms through the organ (21). Filarial worms most often cause a lingering variant of PIE known as tropical pulmonary eosinophilia (TPE) (22-24). TPE occurs in only a small percentage of patients (mostly...
males) with bancroftian filariasis (25,26), and nonfilarial helminths can cause a similar clinical syndrome (27).

Filarial TPE is a syndrome clinically characterized by symptoms of bronchial asthma with paroxysmal nocturnal cough and anorexia. Pulmonary function tests reveal restrictive and obstructive lung disease. Interstitial lung disease is believed to result from immune hyperreactivity to microfilaria. Microfilariae are trapped and destroyed in the lung by antibody-dependent, cell-mediated cytotoxicity involving eosinophils (28). This triggers an inflammatory process that evolves over time. Degenerating worms release somatic allergens that bind to specific cell-bound IgE and thereby trigger the release of vasoactive and inflammatory molecules by lung basophils and mast cells. These mediators cause some of the allergic asthma-like manifestations of TPE. Many empirical and experimental observations are consistent with this model. Microfilariae or recognizable fragments are present in lung nodules from patients with TPE and thus may serve to trigger and focus the pathogenic process (29). Most postulated effector cells and molecules are present in lower respiratory tract epithelial lining fluid, i.e., at the site of the lesions (30,31).

Microfilariae are a primary trigger of lung disease in the pathogenic scheme just described: the continuous arrival in the lungs of new microfilariae produced by living adult worms located elsewhere and identified by ultrasonography (27) provides the stimulus that guarantees the chronicity of the clinical syndrome. However, the effector mechanisms that damage lung parenchyma are not yet completely known. Possibilities include immune complex deposition, eosinophil-mediated injury to parenchymal cells and the interstitial matrix (32), cytokine-mediated cell injury and other so-called bystander effects, as well as possible autoimmune reactions triggered by crossreactivity between filarial and human collagens (33). Alternatively, factors released into the general circulation by filarial worms lodged in extra-pulmonary lymph vessels may act on lung epithelial cells and cause airway hyperreactivity and/or tissue damage (34). The lung is the main organ affected by TPE, but extra-pulmonary lesions such as hepatosplenomegaly and lymphadenopathy can occur (23). Cardiovascular involvement in TPE has been reported (35-38). Because the electrocardiographic changes suggestive of ischemic heart disease are reversed following treatment with DEC, it is assumed that carditis is of filarial etiology. This may not be true, however, because in addition to being an antihelminthic, the drug has anti-inflammatory, antimicrobial, antifungal and antiviral properties (39-42).

TPE is the best known clinical manifestation of occult filariasis (43). Destruction of microfilariae in organs other than the lung can cause several other clinical entities. Bonne (44) reported the presence of microfilariae and hypereosinophilia in the spleen, but not in other internal organs, of a man who died a violent death. The histological changes in the spleen were similar to those described by Meyers and Kouwenaar (23) in lymph nodes from patients with TPE.

The relationship between filarial worms and tropical endomyocardial fibrosis (TEMF) remains unclear. The geographic distribution of TEMF overlaps with that of TPE in India and with loiasis in Africa, suggesting that eosinophils induced and activated by filarial infections may cause the observed endomyocardial changes (45). Eosinophils have two main functions: they can inactivate many of the mediators released by mast cells and thereby dampen tissue reactions associated with IgE-mediated degranulation of mast cells. When activated by antibodies or by complement, they can also damage the larval stages of some helminths and kill a variety of cell types. The damage is inflicted by the deposition of the contents of the eosinophilic granule, especially the major basic
protein (MBP), on the surface of the target
worm or cell. It has been suggested that MBP
release may partly account for the tissue
damage observed in conditions of local
hypereosinophilia such as TEMF. Whether
this has any relevance to the cardiac changes
seen in TPE is unknown, but eosinophils
from patients with TPE are partly degranu-
lated in vivo (46).

**Microfilarial granulomata**

Circumscribed granulomata containing
microfilariae have been observed in the
spleen or occasionally in other organs (47,48).
However, considering the large number of
microfilariae present in most infected indi-
viduals, microfilarial granulomata are ex-
remely uncommon and rarely cause clinical
illness (49).

**Filarial splenomegaly**

The spleen is not usually involved in
bancroftian or brugian filariosis although
splenomegaly occurs in experimentally in-
fected animals. Single-dose DEC treatment
reduced splenomegaly in residents of an area
of Papua New Guinea where bancroftian
filariosis and malaria are co-endemic, a find-
ing consistent with the notion that filarial
infection associated with malaria resulted in
higher spleen rates and sizes (50). This does
not establish a causal relationship between
*W. bancrofti* and splenomegaly because DEC
has many properties besides being an anti-
helminthic (see above). Further, reduction
of splenomegaly in patients with concomi-
tant filariasis and malaria can be achieved by
treatment with antimalarials without associ-
ated DEC (51).

**Skin rashes**

Transient urticarial skin rashes and non-
suppurative swellings without signs of acute
inflammation were observed in World War
II military personnel in the South Pacific
(52,53) and in human volunteers infected
with *B. malayi* (54), but appear to be rare in
native residents (55). Pathological changes
in biopsies from transient rashes in World
War II soldiers are consistent with an aller-
gic reaction (56) and markedly differ from the
inflammatory reactions observed in the
skin of patients with chronic lymphedema
(3). Some consider these fugitive rashes to
be the equivalents of Calabar swellings (52),
i.e., transient areas of angioedema thought to
result from an allergic response to *Loa loa*
products. Of interest, loiasis in temporary
residents also often presents with increased
severity and frequency of pruritus, urticaria
and Calabar swellings compared to the na-
tive population (57). Thus, immune reac-
tions to filarial worms in immigrants qualita-
tively and quantitatively differ from those in
native residents, perhaps because the latter
develop pre- or perinatal immune tolerance
to worm antigens (58,59).

**Final considerations**

Infection with lymphatic filarial para-
sites is associated with a remarkably wide
range of clinical signs, symptoms and se-
quelae. Because adult worms are considered
to be the main parasite stage responsible for
clinical manifestations of the infection, the
importance of extralymphatic disease mani-
festations has been underrated in the past.
The recent recognition that renal damage
(mainly with microscopic hematuria) is pres-
ent in 30% of asymptomatic male microfilar-
emic carriers (14) clearly shows that the
real extralymphatic morbidity might be un-
derestimated worldwide. Hence, as long as
the lymphatic filariasis elimination program
(60) is not completed and microfilariae are
not eliminated from carriers, it is advisable
that additional studies be conducted in order
to improve our knowledge of extralymphatic
disease in endemic areas of lymphatic filari-
asis.
References


